Identification of Gene Expression Changes Associated with the Progression of Retinal Degeneration in the *rd1* Mouse

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Purpose. One approach to gaining insight into the biological pathways contributing to rod and cone photoreceptor death is to identify patterns of gene expression changes. In the present study, a custom retinal microarray was developed to analyze the *rd1* mouse, a well-characterized animal model of human retinal degeneration.

METHODS. A microarray was constructed containing cDNA fragments corresponding to genes known or postulated to be involved in normal retinal function, development, and disease. Gene expression in *rd1* retina was compared with agematched control retinas at three time points: the peak of rod degeneration (postnatal day [P]14), early in cone degeneration (P35), and during cone degeneration (P50). Selected microarray results were confirmed with real-time PCR. The cellular

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distribution of one of the differentially expressed genes, dick-kopf 3 (*Dkk3*), was assessed by in situ hybridization.

RESULTS. At each stage of degeneration, there was only limited overlap of the genes that showed increased expression, suggesting the involvement of temporally distinct molecular pathways. Genes active in transport mechanisms and in signaling pathways were differentially expressed during rod degeneration, whereas genes with functions in protein modification and cellular metabolism were differentially expressed during cone degeneration. Increased expression of genes involved in cell proliferation pathways and oxidative stress was observed at each time point.

Conclusions. These microarray results provide clues to understanding the molecular pathways underlying photoreceptor degeneration and indicate directions for future studies. In addition, comparisons of normal and degenerated retina identified numerous genes and ESTs that are potentially enriched in rod photoreceptors. (*Invest Ophthalmol Vis Sci.* 2004;45: 2929–2942) DOI:10.1167/iovs.03-1184

The molecular events that contribute to retinal degeneration **1** are highly complex, reflecting in part the interdependence of different retinal cell types and the intimate functional and structural contacts between photoreceptors and the adjacent retinal pigment epithelium. Mutations in more than 80 genes, mostly expressed in rod photoreceptors, have been implicated in retinal degenerations, and more than 50 additional disease loci have been mapped (RetNet, http://www.sph.uth.tmc.edu/ Retnet/home.htm; provided in the public domain by the University of Texas Houston Health Science Center, Houston, TX). Many of the disease-causing genes have been identified as key players in retinal activities, including enzymes involved in phototransduction and the visual cycle, transcription factors that control retina-specific gene expression, and structural proteins that support the unique outer segment structure of photoreceptors. 1-3 Despite knowledge of the functions of many of these proteins, the cellular and molecular pathways leading from the primary mutation to cell death are not understood. An additional level of complexity exists for the pathogenesis of the more common retinal diseases, such as age-related macular degeneration (AMD) and glaucoma, which are believed to involve the interplay of multiple genetic and environmental factors.

Although loss of rods has substantial functional consequences such as night blindness and some constriction of visual fields, rod death by itself does not cause severe vision loss. Rather, it is the delayed wave of nonautonomous cone photoreceptor degeneration that results in blindness. The mechanism by which death of rods leads to death of cones is unknown. Several lines of evidence suggest that cone viability may depend on diffusible or contact-mediated factors, ⁴⁻⁶ but it is not yet established whether their absence results in cone

death. It has been proposed that these survival factors may be rod derived.⁷ Identification of gene expression changes that accompany cone degeneration is an important step toward distinguishing between these possibilities and generating new hypotheses.

Naturally occurring and experimentally generated rodent models have been valuable for exploring the histopathological, functional, and molecular changes that occur during retinal degeneration.^{8,9} One well-studied model is the retinal degeneration (rd1) mutant mouse, which has a deficiency in the rod photoreceptor-specific cGMP phosphodiesterase β-subunit (BPDE), a mutation that also causes retinal degeneration in humans. ¹⁰ The phenotype of the *rd1* mutant allows one to address several questions applicable to many of the retinal dystrophies. For example, although early changes in cGMP and calcium dynamics^{11,12} can be directly attributed to the β PDE mutation, the molecular events that lead to rod death and delayed cone degeneration are unclear. Recent data have described morphologic and molecular changes in inner retinal cells during and after photoreceptor degeneration in rd1 and other models, 13-15 yet the role of these cells in promoting or protecting photoreceptors from apoptosis is unknown. In addition, a variety of neurotrophic factors have been reported to reduce the rate of photoreceptor loss in rd1 and other retinal degenerations (Refs. 16, 17 and references therein). Although Müller cells and microglia have been implicated in this process, 18-20 the molecular mechanisms involved remain to be established.

Because rod and cone photoreceptor death and the reactive changes that occur in neighboring retinal cells are likely to involve the coordinated expression of multiple genes, one approach to identifying pathways involved, as well as individual disease-causing genes, is to monitor alterations in gene expression associated with degeneration. 21,22 Among the methods being used to profile retinal gene expression, data mining of cDNA libraries²³⁻²⁵ and serial analysis of gene expression (SAGE)²⁶ are well suited for in-depth quantitation of gene expression under a single experimental state. In comparison, microarray technologies have the advantage of being able to measure gene expression changes across multiple experimental conditions or different disease states. 27-29 In a study of the rd1 mouse utilizing a 588-gene cDNA macroarray, Jones et al.³⁰ found increased expression of one gene, nucleoside diphosphate kinase nm23-M2, which is involved in transcriptional regulation and suppression of metastasis.³⁰ With a 205gene apoptosis-related macroarray, upregulation of frizzledrelated protein-2 (SFRP2) was detected in donor retinas from patients with retinitis pigmentosa,31 and subsequent experiments have provided additional data implicating involvement of the Wnt signaling pathway.³² Further demonstrating the power of these approaches, both SAGE³³ and microarray analysis³⁴ have helped in the identification of inosine monophosphate dehydrogenase type 1 (IMPDH1) as the mutated gene in a form of autosomal dominant retinitis pigmentosa (RP10).

In this study, we developed a custom mouse retinal cDNA microarray that includes genes that have predicted expression in retinal neurons and glia. We used the array to analyze gene expression in degenerating *rd1* retina and identified genes and molecular pathways that are potentially involved in rod photoreceptor death. Furthermore, by comparing a time series of degeneration, we identified gene expression changes that may mediate disease progression—in particular, the non-cell-autonomous death of cone photoreceptors. An additional finding was the identification of novel retina-enriched genes and expressed sequence tags (ESTs), several of which map to known human disease loci and can be considered as potential candidate genes for retinal diseases.

METHODS

Custom Retinal Array Preparation

The mouse retinal microarray has been described in detail elsewhere. 35 The cDNAs on the retinal array represent 5376 genes and were assembled from individually purchased IMAGE clones and from selected clones in a mouse eye cDNA library. The purchased clones were chosen by searching the Online Mendelian Inheritance in Man (OMIM), PubMed, and UniGene EST databases at the National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/; provided in the public domain by the National Institutes of Health, Bethesda, MD)³⁶ for all genes with known roles in normal retinal function, development, and degeneration and genes implicated in retinal diseases. To include genes that may not have been selected by these criteria, we also searched the UniGene database (NCBI; http:// www.ncbi.nlm.nih.gov/UniGene) for all murine ESTs that were expressed in retina and brain and that were obtainable. The selected IMAGE clones (3752 unique sequences) were purchased from Research Genetics (Huntsville, AL). An additional 1624 clones were obtained from a sequenced adult mouse eye cDNA library to increase the gene set, generously provided by Jeremy Nathans and Amir Rattner (The Johns Hopkins University). Functional annotation was performed with the Source and Go databases (http://genome-www5.stanford. edu/cgi-bin/source/sourceSearch and http://geneontology.org/), 37,38 and manual annotation with NCBI PubMed (http://www.ncbi. nlm.nih.gov/PubMed/) and LocusLink databases (http://www.ncbi. nlm.nih.gov/LocusLink/), to confirm or provide missing annotations. Attempts to include the primary or major function of each gene were made. In some cases, more than one function was assigned.

The clones were rearrayed into 96-well plates, PCR amplified, purified, and eluted in TE. PCR-amplified clones were suspended in a final concentration of 50% dimethylsulfoxide (DMSO) and arrayed onto silylated slides (SuperAmine; Telechem, Sunnyvale, CA) using an arrayer (Microgrid II; Biorobotics, Cambridge, UK) with 100- μ m-tip quill pins (Biorobotics). The spots were printed in duplicate on the array, separated by half the length of the slide. Spot quality was verified by fluorescence staining (Sybr-Green II; Molecular Probes, Eugene, OR) on several slides from each print.

RNA Isolation and Probe Labeling

All procedures involving mice were performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research and were approved by the Animal Care and Use Committee at The Johns Hopkins University. Tissue samples were obtained from homozygous rd1/rd1 mice on a C57BL/6 background (for postnatal day [P]14 and P50),³⁹ or homozygous rd1/rd1 mice on a C3H background (for P35) and age-matched control C57BL/6 mice and were processed immediately or flash frozen and stored at -80°C. Retinas were dissected under a microscope to exclude pigmented epithelium and other extraretinal tissue. Total RNA was isolated using phenolchloroform extraction (TRIzol; Invitrogen, Carlsbad, CA). Retinas from four animals were pooled to account for the small amount of retinal tissue and low RNA yield per retina and also to minimize the effect of biological variability. A reference sample was made containing 30% retina RNA and 70% brain RNA, derived from a mix of strains at various ages, from juvenile to adult. RNA integrity was assessed by gel electrophoresis with A₂₆₀/A₂₈₀ absorbance ratios and by analyzing an aliquot (Bioanalyzer; Agilent, Palo Alto, CA).

Microarray Hybridization

Details of arraying and hybridization methodologies are included in the Supplemental Methods (available at www.iovs.org/cgi/content/full/45/9/2929/DC1) and were reported previously.³⁵ Probe preparation and hybridization were based on published protocols.^{40,41} A minimum of four replicates were performed of each experiment, and dye swaps were used to minimize the potential for differential dye effects. The RNA used in replicate experiments for the mutant mice was from

different preparations. A reference sample experimental design (wild-type versus reference and mutant versus reference) was used for time points P14 and P50, and a direct comparison design (wild-type versus mutant) was used at time point P35. This difference in comparison design was not predicted to affect the final results because each time point was analyzed separately to determine significant differential expression, and all subsequent clustering analyses were performed on the calculated mutant-wild-type ratios. Furthermore, we calculated the distribution of the expression ratios at each time point by dividing the ratios into groups that increased by $0.1 \log_2$ units. The distributions for the three time points were very similar, indicating that the microarray results from the two designs are highly comparable and that the mixed design did not introduce experimental bias (see the Results section and Fig. 2B).

Twenty micrograms of total RNA were labeled using an indirect dye-incorporation method with reverse transcriptase (SuperScript II; Invitrogen), random hexamers, and aminoallyl dUTP/dNTP (final concentrations: 1.25 mM dATP, dCTP, and dGTP; 1 mM dTTP; and 0.25 mM aminoallyl dUTP [Sigma-Aldrich, St. Louis, MO]). After removal of unincorporated nucleotides, the DNA was coupled to fluorescent dyes by incubation with monoreactive NHS-cy3 or -cy5 dye (Amersham Biosciences, Inc., Piscataway, NJ) for 2 hours at room temperature. The aminoallyl-labeled cDNA probes were hybridized to the printed slides in 50% formamide, 5× SSC and 0.1% SDS at 42°C overnight in a vacuum-sealed hybridization chamber (GeneMachine, San Carlos, CA). Repetitive sequences were blocked with polyA DNA and Cot fraction mouse DNA (Invitrogen). The hybridized arrays were washed at 42°C for 4 minutes in $1\times$ SSC and 0.2% SDS, followed by two 4-minute room-temperature washes, in 0.1× SSC and 0.2%SDS and then 0.1× SSC. The slides were dried with an air can and fluorescence images of the hybridized microarrays were generated by a scanner and its accompanying software (Scan Array 5000; Perkin Elmer, Boston, MA).

Data Analysis

Spot-finding, image analysis, and quantification were performed on the scanned microarray data TIFF files (Imagene software; Biodiscovery, Inc., Marina del Rey, CA). A linear combination method was applied to calculate the adjusted intensity measurements for arrays with multiple scans, as described in Chowers et al.41 Local background was subtracted from the spot signals, and the data were log₂ transformed. Low-intensity spots were flagged and then treated by flooring to a threshold defined by the average intensity of all low-intensity spots within the same array. Flagged spots with intensity lower than the threshold were upgraded to the threshold, whereas spots with intensity higher than the threshold remained at the original values. This floor method maximized the proportion of informative spots included for later analysis, and at the same time minimized the effect of poorquality data arising from low-intensity signals. If a spot had both cy3 and cy5 channels floored, it was not included in the normalization process, and the final log ratio of this spot was assigned as zero. For all other spots, we first did loess normalization based on multivariate analysis (MVA) plots. 42 Plate effects and pin effects were checked for each array based on box plots. For arrays with a plate/pin effect, a median normalization was applied within each plate/pin, and the median log ratio was subtracted from the loess-normalized log ratio. Statistical analysis of the normalized and transformed data was performed with the Significance Analysis of Microarrays (SAM) program created by Tusher et al.43

An alternative clustering analysis was also developed because our experiments included a limited number of time points and experimental conditions. T statistics were calculated between all successive time points as follows: Suppose there are n time points in the experiment. First, t statistics are calculated for the n time points for each gene t as

$$t(i) = \frac{\bar{x}_{\text{m}}(i) - \bar{x}_{\text{w}}(i)}{\sqrt{\frac{V_{\text{m}}(i) + V_{\text{w}}(i)}{n_1 + n_2 - 2}(1/n_1 + 1/n_2)}}$$

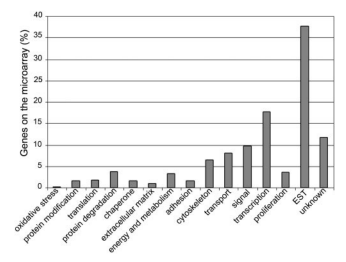


FIGURE 1. Functional annotation of genes on the mouse custom cDNA array. The percentage of genes in each functional class is shown, out of the total number of genes in the array.

where \bar{x}_m and \bar{x}_w are the average expression ratios for mutant and wild-type, V_m and V_w are the within-group variances for mutant and wild-type, and n_1 and n_2 are the number of measurements for mutant and wild-type. Next, considering two successive time points, we defined expression changes of "up," "down," or "no change," by choosing a set of threshold values for the t statistics. With combinations of different expression changes at each time step, all possible expression patterns can be constructed. For example, for an experiment with three time points, we would have eight different patterns ("up-up," "up-down," "no change-up," and so on). Permutation analysis was performed to evaluate the statistical significance for each pattern. For each expression pattern, we counted the numbers of genes that have satisfied the criteria for the pattern before and after the permutation, and the ratio between these two numbers was used as a conservative estimate of the false-discovery rate.

The statistical significance of the distribution of functional classes of the differentially expressed genes was determined as follows: If the differentially expressed genes tend to come from a particular functional group, the frequency of differentially expressed genes for this group would be significantly higher than its corresponding frequency in the total gene set on the array. Binomial and Pearson χ^2 tests⁴⁴ were performed to check whether the distribution of the percentage of differentially expressed genes that fall into various functional categories is different from the distribution of the percentage of all genes that fall into each functional group. When each functional group is considered separately, the number of genes that fall into a single group follows a binomial distribution. Based on the null distribution of genes in the functional classes (Fig. 1), we calculated the probability for the number of observed differentially expressed genes that fall into a functional group.

Quantitative PCR

Total RNA was extracted from retina tissue from a pooled sample of four animals, as described earlier. cDNA was synthesized from 1 μ g total RNA with reverse transcriptase (Thermoscript; Invitrogen). Quantitative real-time PCR (QPCR) was then performed (Lightcycler-Fast-Start DNA Master SYBR Green I kit; Roche Diagnostics, Nutley, NJ), according to the manufacturer's instructions. Each QPCR assay reaction also included amplification of the control genes actin or acidic ribosomal phosphoprotein P_0 (ARP) 45 from each cDNA reaction. Primers were chosen from exons separated by large introns, and the PCR reaction quality and specificity were verified by melting-curve dissociation analysis and gel electrophoresis of the amplified product. For quantification, a standard curve was generated from the cDNA tem-

plate. Relative transcript levels of each gene were calculated using the second derivative maximum values from the linear regression of cycle number versus log concentration of the amplified gene. Amplification of the control genes was used for normalization. Primer sequences are included in the Supplemental Methods available online.

In Situ Hybridization

Probe preparation and hybridization of mouse dickkopf 3 was performed as previously described, with minor modification. ⁴⁶ The detailed protocol can be found in the Supplementary Methods.

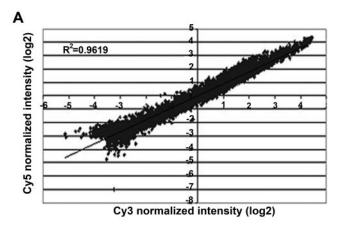
RESULTS

Design and Optimization of the Microarray

The mouse retinal custom array was designed to include genes that would be likely candidates for involvement in normal and disease processes in the retina. Genes were chosen based on function or tissue distribution. The first group included genes involved in essential cellular mechanisms, such as transcription and signaling; genes that had known activities in the retina, such as phototransduction and outer segment structural proteins; and genes implicated directly in disease by identified mutations, or indirectly, as part of general disease processes of neurodegeneration and retinopathies, such as apoptosis and angiogenesis. The genes were selected by searching literature and online database resources (NCBI, PubMed, OMIM; see the Methods section). In addition, we wanted to include all genes expressed in the retina, regardless of whether their function was known, by selecting all available mouse retina ESTs. Because a limited number of mouse retina libraries were in UniGene at the start of the project, we also selected available genes and ESTs expressed in the brain to have a large representation of neuronal and glial genes on the array. We also included 1624 clones from a sequenced adult mouse eye cDNA library (Jeremy Nathans and Amir Rattner, The Johns Hopkins University). In total, we obtained 5376 clones for the murine retina nonredundant gene set. The genes were annotated into the classes shown in Figure 1. Of note, more than one third of the genes in the set included ESTs, predicted genes, and genes of unknown function.

Various optimization and validation studies for probe labeling and hybridization were performed with the printed arrays. For example, in our experience, PCR products resuspended in 50% DMSO produced better spot morphologies and hybridization intensities than those in SSC-containing buffers (3× SSC or 25%DMSO/3× SSC; data not shown). We also found that aminosilane-coated slides produced more consistent results than those that were polylysine-coated, and various aminoallyl UTP-to-TTP ratios were tested to determine which gave the most favorable dye incorporation and signal-to-noise ratios. All subsequent analyses were performed with the optimized protocol described in the Methods section, which is similar to the protocol we recently reported for human retinal arrays. 41

To assess microarray reliability, identical samples were compared using a self-self hybridization. Scatterplot analysis revealed a high degree of correlation, with an R^2 of 0.9619 (Fig. 2A). Only seven genes of the 10,752 spots on the slide (0.065%; each gene printed twice) showed a greater than twofold ratio of normalized cy5/cy3 values. As expected, low-intensity signals were more variable: All the genes that had spurious differential expression had intensity levels close to background. This test also demonstrated that there was minimal artifactual variation from dye effect.



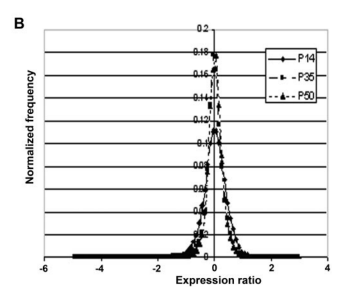


FIGURE 2. Validation of microarray hybridization. (A) A reference sample was hybridized to itself in a dye-swap experiment, demonstrating that there was minimal artifactual variation due to dye bias. Normalized intensity values (\log_2) are shown for cy3 and cy5 scans for each gene on the array. The correlation coefficient (R^2) is indicated. (B) Overlapping distribution of expression ratios at P14, P35, and P50. Expression ratios, in increasing 0.1 \log_2 units, were plotted against normalized frequencies. The overall distributions of expression ratios were very similar at all three time points. The two later time points, P35 and P50, were essentially indistinguishable. The distribution at P14 was slightly wider than that at P35 and P50, indicating that at the P14 time point rd1 and wild-type retinas are more different from each other than they are at the other two points.

Identification of Genes Differentially Expressed during Retinal Degeneration

A time-series experiment was performed to compare the gene expression profiles of normal and degenerating homozygous *rd1* retinas. The time points analyzed (P14, corresponding to the peak of rod degeneration; P35, post-rod and early cone degeneration; and P50, during cone degeneration) were chosen in an effort to maximize the likelihood of identifying genes that promote progression of degeneration, particularly genes that contribute to cone photoreceptor death.

To determine whether experimental bias was introduced by the mixed experimental design used in this study (see the Methods section for the experimental design used) we calculated the distribution of the expression ratios of the three time points (P14, P35, and P50). The overall distributions of expression ratios were very similar at each time point (Fig. 2B), indicating that the results from the pair-wise and reference designs are comparable and that there was no detectable introduced bias. The two later time points, P35 and P50, completely overlap, suggesting that the observed larger number of expression changes at P35 (described later) reflects true biological events and are not an artifact due to the study design.

The SAM algorithm⁴³ was used to determine the likelihood that observed expression level changes reflect true expression level differences ("real" change) rather than artifacts due to chance (see the Methods section for the normalization and statistical protocols used). Instead of using a predefined n-fold ratio cutoff, a false-discovery rate (FDR) was calculated by considering the variability in experimental measurements and comparing the expression ratios from the microarrays with ratios calculated by randomly permuting the control and experimental groups. Gene expression changes were considered significantly different between wild-type and mutant if they were identified in the SAM analysis using a low FDR. We first established which genes were differentially expressed at each of the time points by using FDR levels in the same low range (high stringency: 1.2% for P14, 0.2% for P35, and 0.5% for P50). At P14, 70 genes were differentially expressed: 69 decreased and 1 increased. At P35, 177 genes were decreased and 33 were increased, and at P50, 144 genes were decreased and 62 were increased (Table 1 and Supplemental Table 1 at www.iovs. org/cgi/content/full/45/9/2929/DC1). Increasing the FDR for the P14 analysis to 16% resulted in a larger number of differentially expressed genes (118 decreased and 62 increased). We chose to use this FDR level to allow for comparable subsequent analyses and clustering of the three time points, realizing that a fraction of the identified genes would not represent truly differentially expressed genes.

Confirmation of Differentially Expressed Genes

OPCR was used as an independent method to verify a subset of our results and to assess the reliability of the microarray data. We tested 11 genes in mutant and wild-type retinas at all three time points (33 expression changes in total). The genes were chosen to assess a range of expression ratios, including genes that had high differential expression and those that were unchanged on the arrays. The results of the QPCR analysis are shown in Figure 3 (see also Supplemental Table 2 at www.iovs. org/cgi/content/full/45/9/2929/DC1). There was a high correlation between array and QPCR mutant/wild-type ratios for the 11 genes tested ($R^2 = 0.8736$). The direction of expression differences for mutant versus wild-type was replicated by QPCR with only three exceptions in the 33 changes tested. Furthermore, the QPCR results confirmed the dynamic change in expression observed on the array across the three time points. The absolute ratios differed between QPCR and the array, a finding that has also been reported in other stud-⁻⁴⁹ Therefore, the replication of the temporal pattern of ies.4 gene expression changes by QPCR suggests that the microarray results accurately reflect biological changes.

Temporal Patterns of Differential Expression during Degeneration

Functional group analysis in microarray experiments can help provide insight into the molecular pathways involved in the biological process under study. However, in interpreting such analyses, it is important to exercise caution, because the assignment of genes to particular functional classes can sometimes be difficult and subjective. Some genes have multiple functions, some genes have functions that are only incompletely defined, some functional groups can be overlapping, and our knowledge of gene function is continually changing.

With these caveats in mind, the genes on the murine retinal array were divided into functional classes (Fig. 4) using the approach described in the Methods section. This comparison illustrates the overrepresentation of certain gene classes at each time point compared with their frequency in the gene set, suggesting their functional importance (Fig. 4A, Table 1). Of note is the large representation of genes involved in signaling processes at P14 and P50 compared with the whole array, transport genes at P14 and P35, and metabolism genes at P35 and P50. Genes involved in oxidative stress were most prominent during degeneration at P35 (1.9% vs. 0.2% in the entire gene set). Further subdivision of the differentially expressed genes into groups with increased and decreased expression and comparison to the entire gene set confirmed these trends (Figs. 4B, 4C).

The resultant classification made evident some potentially interesting differences between the P14, P35, and P50 time points. For example, equivalent numbers of protein-modification genes had increased or decreased expression at both P14 and P35, whereas at P50 there were three times more genes involved in protein modification that had increased rather than decreased expression (Figs. 4B, 4C). Similarly, at P35 there were no signaling genes with increased expression but many with decreased expression, in contrast to the P14 and P50 time points, which had similar numbers of signaling genes with increased and decreased expression (Fig. 4B, 4C). Because modulation of a given pathway might be expected to involve gene expression changes in both directions (for example, inhibitors decreased and activators increased), these differences in the directions of expression change may provide clues to the underlying cellular changes occurring at each time point. To investigate this concept further, we compared the nature of the genes that were changed.

As expected, genes known to be differentially expressed in rod photoreceptors had decreased expression at each of the three time points. In contrast, there was little overlap in the identity of the genes with increased expression. Forty-three genes (of 62) that were upregulated at P50 were not upregulated at P14 (Fig. 4B, Table 1 and Supplemental Table 1). Therefore, the array results may help to distinguish rod degeneration from cone degeneration. For example, different intercellular signaling and protein-modification genes were upregulated during cone degeneration at P50 than during rod degeneration at P14. Several calcium-activated proteins were upregulated only at P50 (e.g., calpain, calmodulin). In contrast, genes involved in cellular transport were more highly represented during rod degeneration than during cone degeneration, particularly those involved in modulating ion transport and neuronal transmission, such as Ly6 and NMDA1 subunit zeta 1. These data suggest the involvement of different molecular events in rod and cone death.

We also compared the annotation of the differentially expressed genes to the frequencies of the functional classes on the entire array (Fig. 4A). We performed binomial and Pearson χ^2 tests to determine whether the distribution of the percentage of differentially expressed genes that fall into various functional categories is different from their distribution in the set of all genes on the array (see Fig. 1 and the Methods section). Several functional groups had P < 0.001 (Fig. 4), indicating that they had significantly higher or lower representation than expected (we chose a conservative probability cutoff because multiple groups were being tested).

The large data set of gene expression changes was clustered into smaller groups to identify genes that were behaving similarly, to infer biological information from the gene expression data. Standard clustering methods, such as hierarchical clustering, self-organizing maps or k-means clustering, are based on the distances between pair-wise gene expression profiles (us-

TABLE 1. Differential Expression at P14, P35 and P50 in rd1 Retinas

Gene	Accession No.	Function
Increased Expression		
P14 Sulfated alvoopratain 2 icoform 2/alvotarin	Mm 106244	Call death
Sulfated glycoprotein 2 isoform 2/clusterin Crystallin beta A4	Mm.196344 Mm.40324	Cell death Unknown function
EST	Mm.27363	EST
Secreted phosphoprotein 1	Mm.321	Adhesion
Selenoprotein P	Mm.22699	Transport
cDNA DKFZp586N1922/CLIP-170-related protein	AL117468	Cytoskeleton
Microtubule-associated protein 1A	M83196	Signaling, structural
Neural cell adhesion molecule	Mm.39188	Adhesion
Dickkopf homologue 3	Mm.55143	Wnt signaling inhibitor, putative tumor suppresso
Glutathione S transferase mu 1	Mm.2011	Oxidative stress, metabolism
P35		
EST highly similar to tissue inhibitor metalloproteinase 2	Mm.206505	EST/enzyme inhibition, metabolism
Vimentin	Mm.7	Structural
Transketolase	Mm.154387	Metabolism
Antioxidant protein 2	Mm.6587	Oxidative stress, structural
Dickkopf homologue 3	Mm.55143	Wnt signaling inhibitor, putative tumor suppresso
Solute carrier family 25 Sulfated glycoprotein 2 isoform 2/clusterin	Mm.298	Transport Cell death
	Mm.196344	EST/metabolism
EST similar to <i>H. sapiens</i> KIAA1077 protein/sulfatase 1 EST highly similar to excitatory amino acid transporter	Mm.45563 Mm.206394	Transport
Secreted acidic cysteine-rich glycoprotein	Mm.35439	Cytoskeleton
P50	WIIII. J J + J J	Cytoskeleton
Membrane glycoprotein M6	Mm.178672	Development, proliferation
Calmodulin	Mm.34246	Signaling
EST highly similar to zygin I (FEZ1)	Mm.5264	EST/adhesion, development
Sulfated glycoprotein 2 isoform 2/clusterin	Mm.196344	Cell death, protein modification
Antioxidant protein 2	Mm.6587	Oxidative stress
EST highly similar to chimerin	Mm.41190	EST/signaling
EST	Mm.226753	EST
Protein-1-isoaspartate O-methyltransferase 1	Mm.258431	Protein modification
Profilin II	Mm.20399	Structural
Ras homologue gene family, member J	Mm.27467	Signaling
Decreased Expression		
P14		
Retinal S antigen	Mm.1276	Signaling, vision
Retinitis pigmentosa 1 homologue	Mm.78749	Signaling, vision
Phosducin	Mm.1205	Signaling, vision
Neural retina leucine zipper Guanine nucleotide-binding protein, alpha transducing 1	Mm.20422 Mm.69061	Transcription, vision Signaling, vision
Rhodopsin	Mm.2965	Signaling, vision
X-linked juvenile retinoschisis protein	Mm.41982	Adhesion, vision
Guanine nucleotide-binding protein beta 1	Mm.2344	Signaling, vision
Dopamine receptor 4	Mm.41075	Signaling
Brain type fatty acid binding protein	AF237712	Unknown function
P35		
Guanine nucleotide-binding protein, alpha transducing 1	Mm.69061	Signaling, vision
Aryl hydrocarbon interacting protein like 1	Mm.4452	Signaling, vision
Guanine nucleotide-binding protein beta 1	Mm.2344	Signaling, vision
Retinitis pigmentosa 1 homologue	Mm.78749	Signaling, vision
EST	Mm.1881	EST
Rhodopsin	Mm.2965	Signaling
Phosducin	Mm.1205	Signaling, vision
EST	Mm.29308	EST
Prominin 1	Mm.6250	Unknown function, vision
Retinal S antigen	Mm.1276	Signaling, vision
P50		
Rhodopsin	Mm.2965	Signaling, vision
Guanine nucleotide-binding protein, alpha transducing 1	Mm.69061	Signaling, vision
Heat shock protein 60 kDa	Mm.1777	Protein modified/heat shock
RIRD-1	BG296148	EST
EST weakly similar to CGI 45 protein	AF151803	EST
Retinal S antigen	Mm.1276	Signaling, vision
Aryl hydrocarbon interacting protein-like 1	Mm.4452	Signaling, vision
Phosducin	Mm.1205	Signaling, vision
EST	Mm.271986	EST
EST	Mm.29308	EST

The top 10 genes that were increased and decreased at each time point determined by SAM analysis are shown. Accession numbers and major function(s) of each gene are listed.

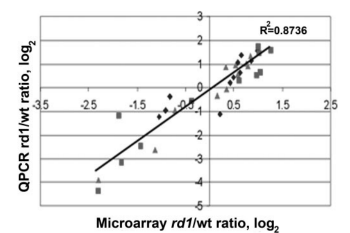


FIGURE 3. QPCR was used as an independent confirmation of the array results for 11 genes (RIRD1 [IMAGE clone 4505626], EST Mm.156168, vimentin, antioxidant protein 2, calcium channel CACNG4, cone-rod homeobox [crx], dickkopf homologue 3, tissue inhibitor of metalloproteinase 2 [TIMP2], Mm.34351, sparc/osteonectin, and Mm.41762). A scatterplot of microarray and QPCR for each gene at each time point shows a high correlation.

ing correlation coefficients). These methods are not optimal if only a few time points or experimental conditions are being compared, as was the case in this study. Therefore, a clustering method was developed that allowed us to determine genes that clustered based on a statistically significant threshold (see the Methods section).

The clustering analysis demonstrated that the temporal expression patterns varied among the genes, Some genes showed continuous increases, others decreased, and others had timepoint-specific expression changes. Because we were interested in the progression of cone degeneration, the genes were clustered based on the change in expression from P14 to P35 and from P35 to P50. Eight different clusters were identified based on the direction of gene expression changes between P35 and P14 and P50 and P35 (Fig. 5; gene lists of each cluster are available in Supplemental Table 3 at www.iovs.org/cgi/ content/full/45/9/2929/DC1). To assess the predictability of this clustering method, we determined which cluster contained known retinal genes. As expected, many of the genes that showed decreased expression at each time point, those in clusters 2 (down-down) and 6 (down-no change), are known retina-enriched genes. The largest number of genes was in cluster 4 (down-up), which contained genes with mutant/ wild-type expression ratios that dipped at P35, and in cluster 3, which had the opposite pattern (up-down), showing a mutant/wild-type ratio that peaked at P35. For example, the expression ratios for follistatin-like 2, insulin-like growth factorbinding protein 7, and several different crystallins peaked during early stages of cone degeneration (up-down), whereas the expression ratios for numerous genes involved in protein turnover, such as proteases, ubiquitin conjugating proteins, and chaperones, were higher during rod and cone degeneration and dipped at P35 (down-up). The coordinated expression changes may indicate that clustered genes participate in similar biological processes.

Dickkopf 3 Localization in Degenerating Retina

Genes with increased expression may be involved in contributing to or inhibiting apoptosis or may be involved in the extensive neural remodeling that occurs after photoreceptor degeneration. To begin to gain some insight into the functional role of the genes that are differentially expressed during degeneration in rd1 mice, an important first step is to determine the cell types in which they are expressed in both the normal and degenerative retina.

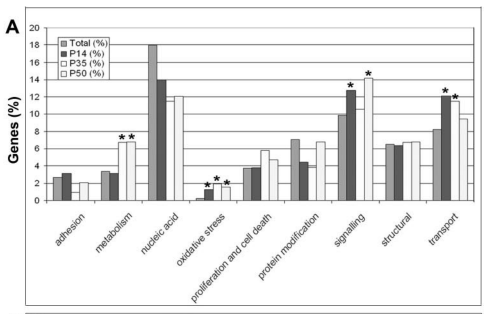
We used in situ hybridization to examine the cellular expression pattern of one of the more interesting differentially expressed genes, dickkopf 3 (Dkk3). Dkk3 is a member of the dickkopf family of Wnt-signaling regulatory genes⁵⁰ and has been implicated in the control of cell proliferation. 51-53 Dkk3 has also been found in a recent study to inhibit Wnt signaling in PC12 cells,⁵⁴ but does not appear to in other contexts.⁵⁵⁻⁵ Expression of the Dkk3 gene has not been reported in degenerating retina, although the expression patterns of several dickkopf family members and Wnt-related genes have been described in the developing and adult normal eye. 58,59 Therefore, we were interested in determining whether the cellular distribution of Dkk3 changes during retinal degeneration.

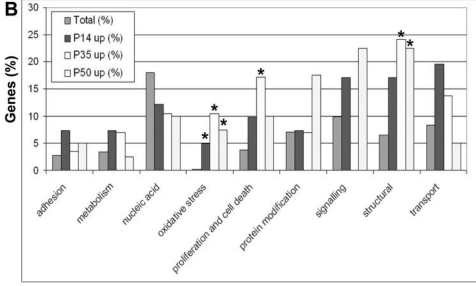
Dkk3 had a more than threefold higher expression in rd1 animals at P35, compared with wild-type mice, and a twofold higher expression at P50. There was 1.6 fold higher expression at P14. Using in situ hybridization, we found that Dkk3 was expressed in inner nuclear layer and ganglion cell layer cells in normal and degenerating tissue (Fig. 6). The overall cellular distribution pattern did not change during degeneration.

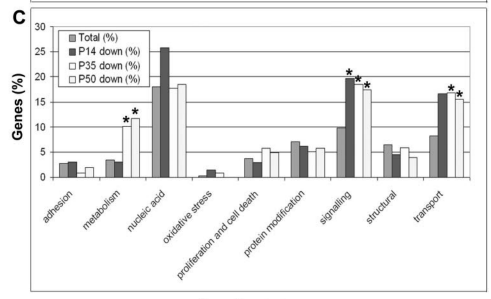
Identification and Chromosomal Localization of Novel Retina Genes

A notable feature of our custom retinal array is the large proportion (over one third) of clones that represent undescribed, or novel, genes, including ESTs and hypothetical and predicted proteins. These clones are an important resource for identifying genes preferentially or specifically expressed in the mouse retina. As might be expected, we found that a significant fraction of the sequences showing decreased expression (16%, 18%, and 20% at P14, P35, and P50, respectively) corresponded to genes already known to be expressed in rod photoreceptors, and many of these have been identified as being mutated in various human hereditary retinal degenerations. Furthermore, many of the ESTs and novel genes were grouped in the down-down cluster along with several known photoreceptor-specific genes, implying that some of the unknown genes may also be expressed in photoreceptors. (It is also possible that some of the genes with decreased expression are expressed in cells in the inner nuclear layer and are differentially regulated during morphologic changes that occur in degenerating retina.) Many of the ESTs with reduced expression in rd1 mice at P35 and P50 may represent genes normally expressed in rod and cone photoreceptors, whereas ESTs with increased expression may represent genes expressed in nonphotoreceptor cells (which have a larger representation in rod-deficient retinas).

The clones on the array (known as well as novel genes) may facilitate the identification of genes that cause hereditary retinal diseases. To generate a list of candidate genes, we determined the chromosomal position of the human counterpart of the genes that had reduced expression in our analyses. As shown in Table 2, 42 of the mouse clones in the down-down and down-no change clusters had human homologues that mapped within chromosomal intervals implicated in retinal diseases (as defined by the RetNet database; known genes with causative mutations, and their corresponding chromosomal regions, were omitted). Despite the large size of some of these regions, the strategy of combining expression data with the chromosome position of the genes and with known retinal disease loci provides a powerful suggestion of a gene's involvement in disease. Curiously, most of the mapped genes are involved in neuronal signaling. Several novel genes were also localized to disease regions and will be interesting to explore further.







Functional class

FIGURE 4. Functional annotation of differentially expressed genes in rd1 retinas during degeneration. (A) The percentage of differentially expressed genes in each functional class at each time point compared with the entire gene set (percentage of total). Based on the null distribution in Figure 1, the probability was calculated for the number of observed differentially expressed genes that fall into a functional group. A functional group with P < 0.001 (*) indicates that this group has significantly higher or lower frequency of genes than expected. (B, C) Percentage of genes with increased (B) and decreased (C) expression in each functional class at each time point compared with the entire gene set (percent of total). A functional group with a P < 0.001 (*) indicates that this group has a significantly higher or lower frequency of genes than expected from the null distribution.

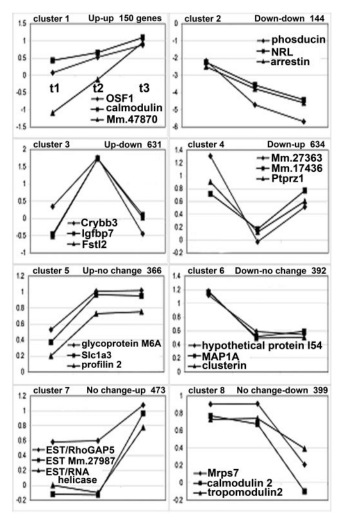


Figure 5. Clustering of genes into eight groups using t statistics. Three genes in each cluster are shown to illustrate the pattern of expression changes. The vertical axis indicates the difference in mutant/wild-type expression ratios between time points P14 (t1), P35 (t2), and P50 (t3). The total number of genes in the cluster is also shown in each graph.

DISCUSSION

In retinitis pigmentosa and other retinal degenerations, mutations in genes predominantly expressed in rod photoreceptors often lead to rod degeneration followed by the death of cones.60-62 To gain insight into the pathogenesis of these diseases, particularly the vision-robbing cone death, a custom mouse microarray was developed and used to characterize retinal gene expression in rd1 mice at specific time points during degeneration. Despite the more rapid degeneration in this animal model compared with most human retinal diseases, rd1 mice offer the benefit of investigating molecular mechanisms that originate from a defined primary mutation that correlates with a particular histologic pattern of cell death. Through this analysis, we have identified groups of genes that are candidate mediators of rod photoreceptor degeneration and may be involved in the initiation and progression of cone photoreceptor death. Future studies will include specific protein quantification of these candidate molecules to determine their significance to photoreceptor degeneration, a necessary step because RNA expression changes do not always reflect protein level changes.

We found particularly provocative the observation that several genes with products known to provide protection against oxidative damage, including antioxidant protein 2,63 glutathione peroxidase 2,64 and paraoxonase,65 were upregulated during the period of cone degeneration. This raises the question, could oxidative damage contribute to cone death after rods disappear? Rods and cones receive their oxygen supply from choroidal vessels that have very high flow and contain fenestrations that allow pooling of plasma, separated from the photoreceptors only by Bruch's membrane and the retinal pigmented epithelium. A high oxygen diffusion gradient allows adequate oxygenation through the entire outer nuclear layer. The elevated oxygen supply is balanced by extreme oxygen utilization by photoreceptors, which are packed with mitochondria to support the activity of the Na/K pump required to maintain the dark current. As rods die, there is decreased oxygen demand, potentially resulting in excessive oxygen supply to the remaining cones. High levels of tissue oxygen increase oxidative stress, which can overwhelm protective mechanisms, as demonstrated by the observation that prolonged hyperoxia causes photoreceptor cell death. 66,67 Thus, it seems reasonable to consider a possible contribution of oxidative damage to cone cell death.

Another interesting observation was that a group of genes usually associated with tissue growth and differentiation showed increased expression in rd1 retina during cone degeneration. Included among this group were the Wnt pathway gene dickkopf 3, membrane glycoprotein M6, insulin-like growth factor binding protein 5, and Bin-1 (also known as myc box dependent interacting protein 1). The products of these genes may act as part of signaling pathways that activate apoptosis, attempt to block apoptosis, or attempt to upregulate protective cell functions. For example, although dickkopf 3 activity in the retina has not been explored, it has been reported to be downregulated in lung carcinoma, 51,52 suggesting that it may participate in the control of cellular proliferation. Increased expression of the Wnt-related genes SFRP and frizzled-4 have been demonstrated in human retinal degenerations, 31,32,68 and Wnt is known to regulate apoptosis in vitro. 69,70 Membrane glycoprotein M6 is associated with calcium-mediated neuronal differentiation of PC12 cells,⁷¹ and insulin-like growth factor-binding proteins are known to mediate neuronal differentiation.⁷²

Genes that had increased expression, particularly genes involved in growth and differentiation, could be part of the glial hypertrophy and proliferation that accompany photoreceptor degeneration. The role of retinal glia cells in cone photoreceptor degeneration in rd1 mice is unclear, although during rod degeneration there are reports of increased glial fibrillary acidic protein (GFAP)⁷³ and immediate-early genes in Müller cells.⁷⁴ In our analysis, we found that vimentin, an intermediate filament protein associated with Müller cell hypertrophy⁷⁵ and apoptosis⁷⁶ was increased at time points P35 and P50. Pleiotrophin, a gene that is found in glia in the brain and is involved in communication with neurons⁷⁷ was increased at P50. Several immune-related genes had increased expression, such as the lymphocyte antigen 6 complex, consistent with activation of microglia,⁷⁸ which have also been implicated in retinal degeneration.²⁰

Müller cells are believed to have a protective effect on rod photoreceptors in several damage paradigms by providing neurotrophins, a well-described class of growth factors. 18,19,79 It is plausible that glial reactive changes attempt to protect cones in rd1, possibly through actions of the growth and differentiation genes that are differentially expressed. Detailed cell localization studies are necessary to demonstrate whether glia cells, and which type, express genes that we found upregulated on the array.

2938

FIGURE 6. Distribution of retinal expression of dickkopf 3 (Dkk3) during degeneration. Microarray and QPCR demonstrated that expression of Dkk3 increased during degeneration. Sections of retina from rd1 and control (C57BL/6) mice were hybridized to probes generated against the antisense strand of the mouse Dkk3 gene. Dkk3 is expressed in cells in the inner nuclear layer and ganglion cell layer; there was no obvious alteration in cellular distribution during degeneration. (A) P14 rd1, (B) P14 BL/6, (C) P35 rd1, (D) P35 BL/6, (E) P50 rd1, and (F) P50 BL/6. The negative control hybridizations, (G) P35 rd1 and (H) P35 BL/6, using a sense probe synthesized against the same region as the antisense strand, did not show detectable hybridization signals. The mutant retina has extensive loss of the photoreceptor outer nuclear layer, clearly evident in P35 and P50. ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer; OS, outer segments; OPL, outer plexiform layer; IPL, inner plexiform layer.

Several crystallin genes also showed expression changes during rod and cone death. For example, crystallin beta A4 was increased 4-fold in mutant P35 retinas and 2.8-fold at P14. A recent proteomics study of rd1 retinas demonstrated that several α - and β -crystallin family members were induced in rod degeneration. ⁸⁰ Similarly, Jones et al. ⁸¹ reported increased expression of α -crystallin at P18 in rd1 retinas, which was dis-

tributed predominantly in the inner retina and possibly in ganglion cells and Müller end processes. In contrast, at P50, when the majority of cones are gone, there was more than a twofold decreased expression of crystallin beta A4 and alpha 1-crystallin.

Crystallins are the major structural proteins in the lens and are present in other tissues, including retina. 82-84 The extra-

TABLE 2. Identification of Candidate Photoreceptor Genes that Map within Retinal-Disease-Critical Regions

Disease and Critical Region	Mouse UniGene	Human UniGene	Gene Name
CORD8 1q12-q24 recessive cone-rod dystrophy	Mm.8	Hs.380869	Upstream transcription factor 1
	Mm.143819	Hs.4788	Nicastrin
	Mm.2261	Hs.148495	Proteasome 26S subunit
	Mm.4598	Hs.6194	Chondroitin sulfate proteoglycan BEHAB/brevican
	Mm.29860	Hs.332012	Immunoglobulin superfamily, member 8
ARMD1 1q25-q31 dominant MD, age-related	Mm.3137	Hs.196384	Prostaglandin-endoperoxide synthase 2
AXPC1 1q31-q32 recessive ataxia, posterior column with RP	Mm.29725 Mm.20429	Hs.75074 Hs.159437	Mitogen-activated protein kinase-activated protein kinase 2 Prospero-related homeobox 1
WILLI KF	Mm.3137	Hs.196384	Prostaglandin-endoperoxide synthase 2
BBS5 2q31 recessive Bardet-Biedl syndrome	Mm.220950	Hs.20999	Hypothetical protein FLJ23142
RP26 2q31-q33 recessive RP	Mm.40678	Hs.41717	Phosphodiesterase 1A, calmodulin-dependent
	Mm.220950	Hs.20999	Hypothetical protein FLJ23142
USH2B 3p24.2-p23 recessive Usher syndrome, type 2;	Mm.29816	Hs.9663	Programmed cell death 6 interacting protein
recessive sensorineural deafness without RP			· ·
WFS2 4q22-q24 recessive Wolfram syndrome;	Mm.33788	Hs.35804	Hect domain and RLD 3
dominant	Mm.3420	Hs.83428	Nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (p105)
MCDR1 6q14-q16.2 dominant MD, North Carolina	Mm.4870	Hs.12477	Synaptosomal-associated protein, 91 kDa homologue
type; dominant progressive bifocal chorioretinal atrophy			
LCA5 6q11-q16 recessive Leber congenital amaurosis	Mm.5135	Hs.301676	Glutamate receptor, ionotropic, kainate 2
1 1	Mm.4870	Hs.12477	Synaptosomal-associated protein, 91 kDa homologue
CYMD 7p21-p15 dominant MD, cystoid	Mm.23567	Hs.82226	Glycoprotein (transmembrane) nmb
AA 11p15 dominant atrophia areata; dominant	Mm.35680	Hs.121619	Chromosome 11 open reading frame 15
chorioretinal degeneration, helicoid	Mm.20837	Hs.20478	Ceroid-lipofuscinosis, neuronal 2, late infantile (Jansky- Bielschowsky disease)
VRNI 11q13 dominant neovascular inflammatory vitreoretinopathy	Mm.205578	Hs.119257	Ems1 sequence (mammary tumor and squamous cell carcinoma-associated (p80/85 src substrate)
• •	Mm.32518	Hs.132004	Cardiotrophin-like cytokine
	Mm.41781	Hs.6059	EGF-containing fibulin-like extracellular matrix protein 2
	Mm.42240	Hs.180677	Splicing factor 1
LCA3 14q24 recessive Leber congenital amaurosis	Mm.46005	Hs.180789	S164 protein
	Mm.5043	Hs.25647	V-fos FBJ murine osteosarcoma viral oncogene homologue
MRST 15q24 recessive retardation, spasticity, and retinal degeneration	Mm.41687	Hs.351327	Hypothetical protein FLJ20452
CORD5 17p13-p12 dominant cone dystrophy, progressive; recessive cone-rod dystrophy	Mm.25722	Hs.41735	Purinergic receptor P2X, ligand-gated ion channel, 1
CACD 17p13 dominant central areolar choroidal dystrophy	Mm.25722	Hs.41735	Purinergic receptor P2X, ligand-gated ion channel, 1
USH1G 17q24-q25 recessive Usher syndrome	Mm.8552	Hs.1578	Baculoviral IAP repeat-containing 5 (survivin)
	Mm.4793	Hs.1437	Glucosidase, alpha; acid (Pompe disease, glycogen storage disease type II)
	Mm.12508	Hs.159557	Karyopherin alpha 2 (RAG cohort 1, importin alpha 1)
	Mm.30176	Hs.50151	Potassium inwardly rectifying channel, subfamily J, member 16
	Mm.21841		Splicing factor, arginine/serine-rich 2
CORD1 18q21.1-q21.3 cone-rod dystrophy; deGrouchy syndrome	Mm.17537	Hs.180933	CpG-binding protein
AIED Xp11.4-q21 Åland island eye disease	Mm.27372	Hs.3383	Upstream regulatory element-binding protein 1
	Mm.92529	Hs.4552	Ubiquilin 2
	Mm.3555	Hs.149155	Voltage-dependent anion channel 1
	Mm.1344 Mm.20873	Hs.765 Hs.211607	GATA-binding protein 1 (globin transcription factor 1) Trinucleotide repeat containing 11 (THR-associated
	M 110/	11- 2055	protein, 230kDa subunit)
DDD Vn11 2 n11 22 natinal dvanlacia maintain	Mm.1104	Hs.2055 Hs.765	Ubiquitin-activating enzyme E1
PRD Xp11.3-p11.23 retinal dysplasia, primary	Mm.1344 Mm.1104	Hs. 2055	GATA-binding protein 1 Ubiquitin-activating enzyme E1
OPA2 Xp11.4-p11.2 X-linked optic atrophy	Mm.27372	Hs.3383	Upstream regulatory element-binding protein 1
The April 1 prinz A mixed optic autopity	Mm.92529	Hs.4552	Ubiquilin 2
	Mm.1344	Hs.765	GATA-binding protein 1
	Mm.1104	Hs.2055	Ubiquitin-activating enzyme E1 (A1S9T and BN75
RP24 Xq26-q27 X-linked RP	Mm.28275	Hs.146381	temperature sensitivity complementing) RNA-binding motif protein, X chromosome

Clones that map within the critical region of various human retinal diseases were identified. Shown for each disease region are the UniGene cluster ID, the mouse gene name, the corresponding human UniGene cluster ID and human gene name. The disease intervals were obtained from the RetNet database.

lenticular function of α -crystallins includes chaperone and antiapoptosis activities, and they play important roles in differentiation, development, and neurodegeneration. 85-87 β -Crystallins have also been reported to have chaperone activity during cellular stress, 86,88,89 but their role in degenerated retina is not known. Crystallins are phosphorylated through cAMP-dependent and -independent pathways, 90,91 suggesting that they interact with signaling-related proteins. α -Crystallin inhibited oxidative stress-induced apoptosis in RPE cells in culture, 92 and in a lens epithelial cell line, oxidative stressinduced apoptosis was associated with decreased α -crystallin. 93 Introduction of α -crystallin into the lens cells prevented H₂O₂-induced apoptosis through inhibition of caspase-3 activation. 93 Based on these studies, it is possible that crystallins are involved in protecting cones from death caused by oxidative damage. This possibility clearly warrants further exploration.

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