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Expression profiling of the developing and mature Nrl^{-/-} mouse retina:

Identification of retinal disease candidates and transcriptional regulatory targets of Nrl

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Abstract

The rod photoreceptor-specific neural retina leucine zipper protein Nrl is essential for rod differentiation and plays a critical role in regulating gene expression. In the mouse retina, rods account for 97% of the photoreceptors; however, in the absence of Nrl (Nrl^{-/-}), no rods are present and a concomitant increase in cones is observed. A functional all-cone mouse retina represents a unique opportunity to investigate, at the molecular level, differences between the two photoreceptor-subtypes. Using mouse GeneChips (Affymetrix), we have generated expression profiles of the wild type and Nrl^{-/-} retina at three time-points representing distinct stages of photoreceptor differentiation. Comparative data analysis revealed 161 differentially expressed genes; of which, 78 exhibited significantly lower and 83 higher expression in the Nrl^{-/-} retina. Hierarchical clustering was utilized to predict the function of these genes in a temporal context. The differentially expressed genes primarily encode proteins associated with signal transduction. transcriptional regulation, intracellular transport, and other processes, which likely correspond to differences between rods and cones and/or retinal remodeling in the absence of rods. A significant number of these genes may serve as candidates for diseases involving rod or cone dysfunction. Chromatin immunoprecipitation (ChIP) assay showed that in addition to the rod phototransduction genes, Nrl might modulate the promoters of many functionally diverse genes *in vivo*. Our studies provide molecular insights into differences between rod and cone function, yield interesting candidates for retinal diseases and assist in identifying transcriptional regulatory targets of Nrl.

Introduction

The mammalian retina contains a diverse array of anatomically and functionally distinct neurons (1). Rod and cone photoreceptors account for over 70% of all cells in the retina. In most mammals, rods are almost 20-fold more in number compared to cones though their distribution may vary greatly in different regions (2). Photoreceptors are highly metabolically active post-mitotic neurons; it is estimated that almost 10 billion opsin molecules are synthesized per second in each human retina (3). Hence, it is not surprising that altered expression or function of opsin and other phototransduction proteins results in photoreceptor degeneration (4-6). The transcriptional regulatory networks underlying photoreceptor differentiation and function are poorly understood.

The neural retina leucine zipper (Nrl) protein, a transcription factor of the Maf-subfamily, is expressed specifically in the rod photoreceptors of the retina (7, 8) and the pineal gland (Mears and Swaroop, unpublished data). Nrl has been shown to interact with the retina-specific homeodomain protein Crx (9) and regulate the expression of rhodopsin (10) and rod cGMP-phosphodiesterase α_{-} (11) and β -subunits (12). In humans, missense mutations of *NRL* are associated with autosomal dominant retinitis pigmentosa (13-17), and in at least one instance (Ser50Thr mutation), the disease may be a result of increased activity of the NRL protein. Targeted deletion of Nrl in mice results in a complete loss of rods and a supernormal S-cone function, as demonstrated by histology, immunocytochemistry, ERG, and expression analysis (18). These observations led to the hypothesis that Nrl plays a critical role in the differentiation of rod photoreceptors, and in its absence, the immature photoreceptors adopt an S-cone phenotype (18). The retina of the Nrl^{-/-} mouse exhibits similarities to the *Nr2e3*^{nd7} mouse (19, 20) and its corresponding human disease enhanced S-cone syndrome (ESCS) (21). One plausible explanation of the phenotypic overlap is that Nrl directly or indirectly regulates Nr2e3 expression, which is undetectable in the Nrl^{-/-} mouse retina (18).

Although several transcription factors have been implicated in photoreceptor differentiation or gene regulation (22-25), their direct impact on the photoreceptor transcriptome has not been elucidated. Microarray-based global expression profiling of tissues from mice deficient in a transcription factor gene can

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point to downstream regulatory targets and provide candidate genes for functional studies and cloning of disease loci (26). This approach has been successfully utilized in studies of the mouse retina (27, 28). The NrI^{-/-} retina is particularly amenable to this analysis because of its dramatic phenotype of no rods and enhanced cones. In the retina of NrI^{-/-} mice, rod bipolar cells have normal morphology, pattern of staining and lamination, and form functional connections with the cones, and the axonal arbors of horizontal cells and AII amacrine cells maintain a normal morphology and stratification pattern (Strettoi, Mears and Swaroop, unpublished data). We, therefore, hypothesize that the comparative analysis of gene profiles from the wild type and NrI^{-/-} retina will, to a large extent, reveal expression differences between rod and cone transcriptomes. Based on our initial analysis of phototransduction genes (18), we predict that transcripts encoding rod photoreceptor proteins would be expressed at lower levels (or undetectable) in the NrI^{-/-} retina. Conversely, the transcripts specific to the normally sparse population of cones are expected to be enriched in the NrI^{-/-} retina.

Here, we report the gene expression profiles, obtained by using Affymetrix GeneChips (MGU74Av2), of wild type and Nrl^{-/-} retina at three time-points (postnatal day 2 and 10, and 2-months). After data normalization by <u>Robust Multichip Average</u> algorithm (RMA) (29) and ranking the statistically-validated genes with a minimum 1.5 average fold-change (AFC) in expression, we have identified 161 differentially expressed genes, which include the known rod or cone-specific genes represented on these Chips. Functional annotation suggests a wide spectrum of physiological changes that likely correspond to differences between rods and cones and/or remodeling of retina in the absence of rods. Our analysis suggests that approximately 25% of all differentially expressed genes identified in this study are either associated with (fifteen) or are candidates (twenty-six) for retinal diseases. Using chromatin immunoprecipitation (ChIP) analysis, a significant proportion of the top ranked genes showing reduced expression in the Nrl^{-/-} retina are demonstrated to be putative direct targets of Nrl, indicating the breadth of its influence on the rod transcriptome.

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Results

Identification of differentially expressed genes in the Nrl^{-/-} retina

The three time-points, P2, P10 and 2-months, were chosen to cover distinct critical stages of photoreceptor development in mouse. In the wild type retina, at P2 many retinal progenitor cells are still exiting the cell cycle and a majority of these will become rods (30). The photopigment of the rod photoreceptors, rhodopsin, is first detected at P4. At P10, retinogenesis is complete, the cells are undergoing terminal differentiation, and photoreceptor outer segments are beginning to form. We chose 2-months of age as another suitable time-point as the retina is structurally and functionally mature and yet old enough to avoid any potential delayed differentiation effects which may occur due to the re-specification of the photoreceptor cell types in the Nrl^{-/-} retina.

To facilitate statistical analysis and identification of "true" positives, four replicate MGU74Av2 GeneChips were utilized for each time-point and strain. Based on absent / present calls (MAS5 analysis), approximately 60% of the probesets (out of $\sim 12,400$) were reported as present or marginally detectable in at least one of the 12 wild type GeneChips, consistent with other studies analyzing single tissue types. Signal quantification and normalization was performed using RMA, a reliable and effective algorithm in control studies (31, 32). The normalized data was then analyzed with a robust two-step procedure to identify statistically significant differentially expressed genes. Due to the tendency of microarrays to quantitatively underestimate fold-change in expression and since RMA normalization compresses the signals (and resulting ratios), an empirical 1.5 AFC cut off was selected as the minimum fold-change (minfc) for statistical analysis. Using these criteria and after removal of those scored as absent on all 24 Chips, a total of 173 probesets were reported as differentially expressed for at least one of the three time-points (i.e., p-value < 1). Of these, 86 show decreased (**Table 1a**) and 87 increased (**Table 1b**) expression in the $Nrl^{-/-}$ retina. The differentially expressed genes are ranked based on increasing false discovery rate confidence interval (FDRCI) p-values, which are similar to FDR p-values except that they account for a specified minimum fold-change level in addition to a level of statistical significance. Although the highly differentially expressed genes are near the

top of the lists as expected, the order is based on both the AFC and the variability of the signal data across the GeneChips. For this reason, probesets displaying a relatively high AFC for a given time-point may still be reported as non-significant (e.g., Nt5e and Fin15 at P10 in **Table 1a** and **1b**, respectively). After removing probesets that belong to the same gene and show similar gene expression profile, a non-redundant set comprising of 78 down-regulated and 83 up-regulated genes is obtained. Almost 90% of these genes are categorized as "known," whereas 18 are novel sequences that are currently represented only in expressed sequence tag (EST) or genomic sequence databases.

Validation by quantitative real-time PCR

Fifty-four different gene/time-point values spanning a broad spectrum of AFC and FDRCI rankings were examined by quantitative real-time PCR (Q-PCR) (**Table 2**). There is a good correlation (R^2 =0.91, data not shown) between AFC reported by microarray and by Q-PCR. Underestimation of the relative degree of fold-change in microarray data is likely due to background noise and limited sensitivity that restricts the dynamic range of this hybridization-based technique. Only three genes (Gas5, Sox11 and 1110002B05Rik) showed disagreement between the two methods (94% validation rate). The discrepancy could be due to the existence of multiple isoforms, which have been identified for these genes. The importance of validation is evident, not just for identifying possible false positives but also to determine the relative fold-change in transcripts (i.e., biological change) compared to the AFC reported by microarray. For example, Guca1a and Kibra are both predicted to be moderately up-regulated (5.6 and 6.0, respectively) in the Nrl^{-/-} retina; however, Guca1a is shown to be up-regulated 5.5-fold by Q-PCR (same as microarray) but Kibra 26-fold (five-fold underestimate by microarray). Similar examples are evident amongst the down-regulated genes. Q-PCR analysis using additional retinal samples for six of the genes revealed similar AFCs (data not shown).

Hierarchical clustering and functional annotation

Relative expression profiling across multiple developmental time-points can provide information on the potential role of a given gene in the context of known biological events occurring within that time frame. Comparison of relative profiles can allow clustering of genes into groups that show similar patterns of behavior. To compare expression patterns between all 161 differentially expressed genes, the average signals from the four replicate GeneChips were first normalized to z-scores, and then run through a hierarchical clustering algorithm. Ten major clusters were identified by visual inspection, and Gene Ontology was used to assign functional annotation of 101 genes (62%) (**Figure 1**).

Cluster I contains genes that display a bimodal (peaks at P2 and 2-month) or constant pattern of expression in the wild type, but show significantly decreased expression at P10 or 2-months of age in the Nrl^{-/-} retina. Cluster II contains three γ -crystallin genes (E, D and F); for these, the peak expression is in the wild type adult retina, but in the Nrl^{-/-} retina there is increased expression at P2 and P10 but a significant decrease at 2-months. Although these genes show AFCs > 2-fold at P10, none of these are considered statistically significant (see **Table 1**). This may be due to the signal noise associated with the high degree of sequence identity between different crystallins. Q-PCR confirmed the decreased expression of Crygd and Crygf at 2-months (see **Table 2**). Crystallins are expressed in neural retina and may play a role in stress response (33). For the genes of Cluster V; their expression peaks at P10 but then decreases (though still detectable) in the wild type adult retina. In the Nrl^{-/-} retina, the peak expression may still be at P10, but is reduced for all these genes, suggesting a potential role in differentiation, as indicated for Ndr1, Ndrl and Lmo1. Cluster VI contains only two ESTs that are expressed across all three time-points but are down-regulated in the Nrl^{-/-} retina.

Almost 80% of genes showing decreased expression in the $Nrl^{-/-}$ retina belong to clusters III and IV. These genes demonstrate an increasing (relative) level of expression, reaching peak expression by P10 (cluster III) or 2-months (cluster IV) in the wild type, suggesting a role in the mature retina / photoreceptors. In the $Nrl^{-/-}$ retina, these genes are down-regulated showing, typically, only a moderate (or no) increase in expression at later time-points. Genes of these clusters are strong candidates for direct positive regulation by Nrl and include Rho, Pde6b and Pde6a (known targets of Nrl) as well as Gnat1 and Gnb1.

The genes showing higher expression in the NrI^{-/-} retina can be organized into four major clusters. The genes of the largest cluster VIII show an increase in expression at P10 or 2-months in the NrI^{-/-} retina relative to the wild type. As anticipated, this includes genes encoding proteins with a role in cone-mediated visual function (e.g., Opn1sw and Gnat2). Expression of cluster VII genes peaks at P2 in the wild type, suggesting a primary role in early development, but in the NrI^{-/-} retina they show elevated expression peaking at P10 or 2-months. Their sustained high expression in the adult retina may be indicative of an aberrant reactivation of gene expression, possibly related to stress, cell death or reactive gliosis. Cluster IX genes show an elevated differential expression (and peak) in the NrI^{-/-} retina, primarily at P10, and may play a role in cone differentiation. Of the 14 genes in this cluster, six are associated with signaling, development or cell cycle/growth. Cluster X includes genes showing peak expression at P2 in the wild type but the expression declines (often rapidly) by P10 or 2-months, suggesting a primary role in early development. In the NrI^{-/-} retina, the expression profile is similar but the expression is elevated and maintained for a longer period.

Direct targets of Nrl, identified by chromatin immunoprecipitation (ChIP)

We hypothesized that targets of Nrl will be enriched among the genes exhibiting reduced expression in the Nrl^{-/-} retina. Hence, we examined the enrichment of the promoter regions that include a potential AP-1 like or Nrl-response element (NRE) of "candidate Nrl targets" by ChIP with a polyclonal anti-Nrl antibody (8) using wild type mouse retina. Twenty different gene promoters were assayed by PCR amplification; of these, eighteen (90%) showed enrichment in the antibody fractions (Nrl-ChIP) over the no antibody control (**Figure 3**), demonstrating *in vivo* promoter occupancy by Nrl. The positive target promoters included three genes (Rho, Pde6b and Pde6a) that are modulated by Nrl. The promoters of other photoreceptor genes (such as Cnga1, Gnat1, Gnb1, Rom1 and Pdc) were also enriched. In addition, a few widely expressed genes, such as Aqp1 (water channel) and adiponectin receptor 1 (AdipoR1), appear to be the target of Nrl regulation in

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mature rods. It should be noted that although only down-regulated genes were analyzed by ChIP, Nrl might negatively regulate (i.e., repress) the expression of cone-specific genes, much akin to the predicted role of Nr2e3 (21).

Identification of retinal disease candidate genes

Many genes showing photoreceptor-enriched expression are associated with retinal disease; these encode diverse functions, including phototransduction (e.g., rhodopsin), transcriptional regulation (e.g., Crx and Nrl), outer segment structure (e.g., Rom1 and Prom1), or maintenance of the extracellular matrix (e.g., Rs1h). Expression profiling of a mouse model with retinal degeneration (Rho^{-/-}) was previously utilized to identify a retinitis pigmentosa disease gene (RP10), inosine monophosphate dehydrogenase type 1 (IMPDH1) (34), which was not an obvious candidate due to its ubiquitous expression and role in guanosine nucleotide biosynthesis. We, therefore, determined the chromosomal location of genes that are differentially expressed in the Nrl^{-/-} retina using *in silico* methods. Based on the map position of the human homolog, 41 of the differentially expressed genes (25%) have previously been associated with or are candidates for retinal diseases (**Table 3**). A few of these (e.g., Mef2c, Nt5e and Cdr2) were also identified in the Rho^{-/-} gene profiling study (34), providing further evidence of their rod-preferred expression. Upregulated genes that are candidates for macular or cone associated diseases include S100A6, RXRG, ADCY2, NP and SOCS3, whereas down-regulated genes that map to the region of rod associated disease loci (such as RP) include NT5E and CDR2.

Analysis of differentially expressed genes

Light response and vision

The genes displaying restricted expression to rods or cones show the most dramatic changes in expression. For the rods, these include genes encoding rod-specific phototransduction proteins such as rhodopsin (Rho), cGMP phosphodiesterase subunits (Pde6a and b), rod transducin subunits (Gnat1 and Gnb1)

and the cyclic nucleotide gated channel subunit (Cnga1). By Q-PCR, transcripts of these genes are virtually undetectable in the NrI^{-/-} retina with expression typically less than 1% of wild type. Modest expression of Pde6a (~7%) and Pde6b (~2%) in the adult NrI^{-/-} retina can be attributed to their expression in non-photoreceptor neurons, as observed for Pde6a (35). Genes encoding cone phototransduction proteins, such as the photopigment S-opsin (Opn1sw), cone transducin subunits (Gnat2, Gnb3 and Gngt2), and the cyclic nucleotide gated channel subunit (Cnga3), show dramatically higher expression in the NrI^{-/-} retina. A number of genes that are expressed in both photoreceptor subtypes show varying degrees of expression change, which may reflect a moderate quantitative bias towards one class (or expression in multiple cell types). These include guanylate cyclase activator 1a (Guca1a or Gcap1), recoverin (Rcvrn), prominin 1 (Prom1), phosducin (Pdc), retinal S-antigen (Sag), retinal outer segment membrane protein (Rom1) and an ATP-binding cassette (ABC) transporter (Abca4). Guca1a displays a 5.5-fold increase in expression in the NrI^{-/-} retina suggesting preferential expression in cones. Notably, although expressed in both rods and cones, mutations in this gene are primarily associated with cone or cone-rod dystrophies (36, 37). Other down-regulated genes may indicate their preferential expression in rods.

Gene regulation, differentiation and development

Transcription factors and signaling molecules that are differentially expressed in the NrI^{-/-} retina may provide insights into the regulatory networks associated with photoreceptor development and/or function. Q-PCR analysis of E14 to P21 retina for the cone photopigment Opn1sw (S-opsin) showed that the increase in its expression occurred at P6.5 in the NrI^{-/-} retina (**Figure 4**). This second-wave of cone differentiation likely corresponds to the post-mitotic photoreceptors that are normally destined to become rods. Therefore, it is predicted that genes associated with rod or cone differentiation would be down or up-regulated, respectively, at this time-point. The expression of MADS-box containing myocyte enhancer factor 2c (Mef2c) (38, 39) is reduced in the mature NrI^{-/-} retina to 20% of the wild type levels. Zfp36l2 (a C3H-type zinc finger protein) is down-regulated approximately 8-fold in the NrI^{-/-} retina. A significant decrease in expression of LIM domain only 1 (Lmo1), a developmentally-associated transcription factor, is observed in the adult retina (10-fold by Q-PCR) suggesting its role in mature rods. Similar profiles are also observed for N-myc downstream regulated 1 (Ndr1) and Ndr-like (Ndr1).

A number of genes encoding transcription regulatory proteins are up-regulated in the Nrl^{-/-} retina. Retinoid X receptor gamma (Rxrg), localized to cones in the adult retina (40) and shown to be induced by retinoic acid (RA) (41), shows 9-fold higher expression in the Nrl^{-/-} retina. Rxrg maps to the region of conedystrophy locus CORD8 (**Table 3**) and is an excellent candidate for this disease. Sal-like 3 (Sall3), a C2H2 zinc finger transcription factor, is required for terminal differentiation of photoreceptors in *Drosophila* (42); its augmented expression is therefore of considerable interest. Validation by Q-PCR, which detects two of the six alternative transcripts, reveals that Sall3 is highly differentially expressed at P10 (20-fold) but is only moderately increased at 2-months (2-fold), suggesting a potential role in cone differentiation. Engrailed-2 (En2), a homeobox transcription factor, shows sustained expression in the mature wild type retina but in Nrl^{-/-} retina it is highly elevated (30-fold increase). The positive regulatory domain zinc finger protein, Prdm1, shows elevated expression (8-fold) in the mature Nrl^{-/-} retina. It is expressed earlier in the wild type retina and is undetectable in the adult.

Apoptosis and stress response

Several genes encoding proteins associated with stress response or apoptosis exhibit decreased expression in the Nrl^{-/-} retina; these include the chaperone heat shock proteins Hsp70.3 (Hspa1a) and Hsp70.1 (Hspa1b). Serum/glucocorticoid regulated kinase (Sgk), which shows peak expression in the adult retina and is down-regulated in the Nrl^{-/-} retina, is shown to be anti-apoptotic and induced in response to multiple forms of stress in epithelial cells (43). Tumor necrosis factor alpha induced protein 3 (Tnfaip3), which inhibits NF-kappa B (Nfkb1) (44), has been associated with light-induced photoreceptor degeneration (45). Tnfaip3 is first detected at P10, and its expression peaks at 2-months. In contrast, Nfkb1 expression is relatively constant in the wild type retina but exhibits a moderate peak at P2. In the Nrl^{-/-} retina, Tnfaip3 is down-regulated 8-

fold, whereas its inhibitory target, Nfkb1, is up-regulated. This observation, may at least in part, provide clues to the mechanism through which stress response and cell death may be mediated in the $Nrl^{-/-}$ retina during late stages (unpublished data). Caspase-7, which is detected in the wild type retina during development, is the only caspase showing elevated (10-fold) expression in the adult $Nrl^{-/-}$ retina.

Calcium homeostasis and retinal function

During the recovery of light response in photoreceptors, cGMP is regulated by cytoplasmic Ca²⁺ via Guca1a (or Gcap1). Both Guca1a and rod arrestin (Sag) are associated with retinal diseases and are differentially expressed in the NrI^{-/-} retina. Calcium/calmodulin-dependent kinase II beta (Camk2b) is up-regulated (15-fold) in the NrI^{-/-} retina. Calcyclin (S100a6) is highly expressed in neurons (46) and shows elevated levels in the NrI^{-/-} retina. The human homolog of this gene maps to a cone-rod dystrophy locus (CORD8). S100a6 is regulated by NF-kappaB (47), which is also augmented in the NrI^{-/-} retina. Two calcium channels genes Trpc1 and Cacnb2 are down-regulated in the NrI^{-/-} retina. Syntrophin acidic 1 (Snta1) is a component of the dystrophin glycoprotein complex (DGC) which may play a significant structural and signaling role (neurotransmission) in the retina (48). Mutations of dystrophin or disruption of the DGC may account for scotopic (rod response) defects in patients with Duchenne muscular dystrophy (49), consistent with rod-enrichment of Snta1 and its down-regulation in the NrI^{-/-} retina.

Melatonin Signaling

Retinal melatonin, acts as a local neuromodulator through the melatonin receptors, which then may control the release of dopamine (50). Three genes of the melatonin pathway, tryptophan hydroxylase (Tph1), dopamine receptor 4 (Drd4) and melatonin receptor 1a (Mtnr1a), are differentially expressed in the Nrl^{-/-} retina. Tph1 is the first enzyme in the biosynthetic pathways of melatonin in the photoreceptors and is believed to be synthesized primarily in the cones (51), consistent with its up-regulation in the Nrl^{-/-} retina. The melatonin receptor 1a, which normally shows peak expression around P2 to P4, is highly elevated in the

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 $Nrl^{-/-}$ retina, and peaks at P8 before rapidly decreasing in expression. The dopamine receptor Drd4, which plays a role in regulating cAMP metabolism, is not highly expressed until P10-P12 in the wild type retina (52), but is down-regulated to less than 10% of wild type levels in the P10 $Nrl^{-/-}$ retina, indicating a role in rods.

Novel functions and novel genes

Although a majority of the differentially expressed genes have a defined function, in many cases their specific role in the retina or their possible bias towards rods or cones is not understood. Deleted in polyposis 1-like 1 (Dp111) is the top FDRCI ranked down-regulated gene and is expressed at <3% of wild type levels. It shows peak expression in the adult retina and is detected in the outer nuclear layer (data not shown) but its function is unknown. A function can be inferred but is not known for calcium activated chloride channel 3 (Clca3), which is up-regulated 44-fold in the Nrl^{-/-} retina. Kibra is a novel WW-domain containing protein expressed primarily in brain and kidney (53) and is up-regulated 26-fold in the Nrl^{-/-} retina. In addition, 18 of the differentially expressed genes identified by microarray analysis match only ESTs. These novel genes could provide new leads for elucidating retinal development and function.

Discussion

Expression profiling and data mining

Appropriate microarray design and data analysis are essential for extracting meaningful results in genome-wide expression profiling studies (54). We utilized RMA for normalization (29, 32) and chose an AFC cut-off of 1.5. A new two-stage gene filtering procedure (55) was applied that controls both false discovery rate (FDR) and minimum fold-change levels. This procedure is based on construction of a set of simultaneous FDR confidence intervals (FDRCI) on the temporal fold-changes of each gene. Genes having at least one confidence interval that covers a range of fold-changes larger than the specified AFC cut-off, which we call minfc, are declared significant at the specified FDRCI level. As FDRCI is more stringent than FDR, the associated significance levels are generally not as high as those of the FDR procedure. For each minfc level studied, the two-stage procedure was used to generate a list of genes ranked according to decreasing FDRCI significance or, equivalently, increasing FDRCI p-value. For an AFC cut-off of 1.5, the complete ranked list, excluding probesets having FDRCI p-values greater than 0.99, consisted of 173 probesets. Of the 54 data points tested by Q-PCR, 51 (94%) were verified. If the minfc is reduced to 1.25, the probeset list is expanded to over 300 probesets (see supplementary Table A). These additional genes may display a reduced validation rate by Q-PCR but add to cluster analysis and pathway construction based on the microarray data. Replicate experiments and statistical analysis are critical for extracting such probesets.

Temporal profiling and clustering analysis add a new dimension to predicting the functional role, possible interactions and regulatory relationships that may exist amongst the genes that are being analyzed. Our studies should identify the genes that are presumably associated with photoreceptor development (P2), terminal differentiation (P10), and function (2-month). Although our data is based on a mixed cell population (whole retina), the generated profiles are dominated by photoreceptors (about 70% of total cells) and can direct future studies to prioritize candidate genes of interest for positional cloning or functional analysis. Of particular interest are the differentially expressed genes encoding proteins associated with visual process, transcriptional regulation, signal transduction and development, as they may provide insights into the regulatory networks and signaling pathways underlying the differences between rods and cones.

Genes encoding metabolism-related proteins represented the single largest class of differentially expressed genes (24%) in the Nrl^{-/-} retina. In addition, one-third of the genes are associated with light response/vision (11%), signaling (18%), and transcription (6%). There was no significant difference between up and down-regulated genes in terms of the specific biological processes affected; however, more genes associated with vision or cell adhesion are down-regulated in the Nrl^{-/-} retina (**Figure 2**). This can be attributed to greater representation of rod- rather than cone-specific transcripts on the MGU74Av2 Chips. A decreased expression of genes encoding structural proteins may reflect the abnormalities of the retinal organization in the Nrl^{-/-} mouse. It should be noted that cones contain more mitochondria as compared to rods (56, 57); expression changes in mitochondria associated genes (Aqp1, Mscs, Skd3 and Clic4) may therefore reflect numerical and physiological differences between the populations of mitochondria in the two classes of photoreceptors.

Expanding the data set: MOE430 GeneChips and custom cDNA arrays

The MGU74Av2 GeneChip contains over 12,000 known genes and ESTs but the retina-specific transcripts are poorly represented. For example, neither Nr2e3 or cone arrestin are on these arrays. Affymetrix has since significantly improved the mouse arrays and the new MOE430 GeneChips now comprise of over 36,000 genes and ESTs. These arrays are superior in design showing greater sensitivity and improved specificity of probesets. One problem with GeneChips is that the probesets are based on public databases and if transcripts are exclusively or predominantly expressed in the retina, they may not have been identified. Custom retinal cDNA arrays (28, 58-60) should therefore complement GeneChip-based analysis of the Nrl^{-/-} retina (Yu and Swaroop, unpublished data).

Differential expression and reactive gliosis in the $Nrl^{-/-}$ retina

The ready-extraction of rod or cone-specific genes from the microarray analysis is complicated by the fact that the $NrI^{-/-}$ retina undergoes a slow form of retinal degeneration (after 4-6 months, unpublished data). A marker of retinal stress, glial fibrillary acidic protein (Gfap), is up-regulated in the $NrI^{-/-}$ retina (18). Reactive gliosis or glial hypertrophy is observed as part of the complex neuronal remodeling that occurs during retinal degeneration (61, 62). Discrimination between photoreceptor-based differential expression and changes due to retinal remodeling must be carefully evaluated, especially when dealing with genes that encode proteins with a poorly defined function. One experimental strategy would be to compare gene profiles, reported here, to those of mouse models of retinal degeneration.

Cones or "cods"

In the original characterization of the Nrl^{-/-} mouse, the photoreceptor population was referred to as "cods" as there was uncertainty as to whether the later developing but functional cones were in fact a type of hybrid photoreceptor. Subsequent analysis with cone-specific markers (such as PNA), suction electrode recordings of isolated photoreceptors (Nikonov, Daniele, Mears, Swaroop and Pugh, unpublished data) and ERG of whole retina, nuclear morphology of the ONL (punctate staining typical of cones) and extensive molecular studies are all consistent with these photoreceptors being cones. Histologically, the retina is abnormal with rosettes and whorls disrupting the ONL, and short, sparse and disorganized OS. These changes, however, may be a consequence of inappropriate nuclear and OS packing within ONL and the sub-retinal region, and may be secondary to the actual identity and differentiation of the photoreceptors. The gene profiling data, presented here, provides strong evidence in favor of the photoreceptors of the Nrl^{-/-} retina

Photoreceptor plasticity and identity

In the absence of Nrl, the failure of the retinal photoreceptors to adopt their appropriate rod identity results in their transformation into cones primarily expressing S-opsin (S-cones). Nrl therefore appears to act

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as a molecular switch during photoreceptor differentiation by promoting the rod differentiation program while simultaneously repressing the cone identity. The suppression of the cone fate is achieved, at least in part, through direct or indirect regulation of the transcription factor Nr2e3 (20, 21), whose expression is undetectable in the Nrl^{-/-} retina (18).

How does Nrl orchestrate the coordinated expression of a broad array of genes that are required for making a mature and functional rod? Delineation of direct downstream targets is the essential first step towards assembling the Nrl-mediated transcriptional regulatory network(s) underlying rod differentiation. Our study has identified several potential direct targets of Nrl by a combined approach of microarray profiling and ChIP. Several of these are known or putative transcription factors or signaling proteins that may play a role in rod or cone differentiation. Comparative retinal gene profiling studies of mouse loss-of-function mutants of other photoreceptor transcription factors (e.g., Crx, Trβ2, Nr2e3) should provide considerable insights into the gene regulatory networks that govern differentiation and homeostasis.

Materials and Methods

Animal Use and Tissue Collection

University Committee on Use and Care of Animals (UCUCA) of the University of Michigan approved all procedures involving mice. Both the $NrI^{-/-}$ mice and wild type controls were of a matched mixed genetic background (R1 and C57BL/6 strains) (18). Mice were sacrificed by cervical dislocation, and the retinas were rapidly excised, frozen on dry ice and stored at -80°C. No signs of pathology were detected in any of the animals used. To isolate sufficient total RNA for labeling protocols, retinas from 2 mice were pooled into a single sample. To minimize false positives due to biological variation, different samples were utilized for four replicate experiments per genotype / time-point (biological replicates). For the developmental Q-PCR studies, retinas were dissected from the embryos of timed-pregnant $NrI^{-/-}$ or wild type females and pooled. Retinas from postnatal time-points were also pooled (entire litter) after dissection.

RNA Preparation

Tissues were placed into TRIzol (Invitrogen, Carlsbad, CA) (added to the frozen tissues at approximately 1.3 ml per 4 retinas) and homogenized (Polytron, Kinematica, Lucerne, Switzerland) at maximum speed for 120 sec. Subsequent steps were done according to the manufacturer's instructions.

Gene Expression Analysis

The GeneChips (Affymetrix, Santa Clara, CA) used in the study contained approximately 12,000 probe sets, corresponding to over 6,000 genes and 6,000 expressed sequence tags (Murine Genome U74A Array v2).

Total retinal RNA was used to generate double-stranded cDNA (ds-cDNA) with SuperScript Choice System (Invitrogen, Carlsbad, CA) and oligo-dT primer containing a T7 RNA polymerase promoter. After second-strand synthesis, the reaction mixture was extracted with phenol-chloroform-isoamyl alcohol, and dscDNA was recovered by ethanol precipitation. *In vitro* transcription was performed by using a RNA

transcription labeling kit (Enzo) with 10 µl of ds-cDNA template in the presence of a mixture of unlabeled ATP, CTP, GTP, and UTP and biotin-labeled CTP and UTP [bio-11-CTP and bio-16-UTP (Enzo Life Sciences, Farmingdale, NY)]. Biotin-labeled cRNA was purified by using an RNeasy affinity column (Oiagen, Valencia, CA), and fragmented randomly to sizes ranging from 35 to 200 bases by incubating at 94°C for 35 min. The hybridization solutions contained 100 mM MES, 1 M NaCl, 20 mM EDTA, and 0.01% Tween 20. The final concentration of fragmented cRNA was $0.05 \mu g / \mu l$ in the hybridization solution. After hybridization, the solutions were removed and GeneChips were washed and stained with streptavidinphycoerythrin. GeneChips were read at a resolution of 6 µm with a Hewlett-Packard GeneArray Scanner. Initial data preparation (i.e., generation of .CHP files) was performed by Affymetrix MICROARRAY SUITE v5.0. Normalization (quantile method) and calculation of signal intensities was performed with the software package RMA from the R project (http://www.r-project.org/). Data was based on four Affymetrix MGU74Av2 GeneChips (biological replicates) for each time-point per genotype (i.e., total of eight GeneChips per timepoint). Of the total 24 GeneChips, only one had to be repeated due to a negative quality report based on raw image and MAS5 analysis. Ratios of average signal intensity (log₂) were then calculated for the probesets (Nrl^{-/-} relative to wild type) and then converted to an average fold change (AFC). Statistical validation was performed on probesets showing a minimum AFC of 1.5. If due to low signal, any of these probesets were reported as having an absent signal (based on MAS5) in all GeneChips (i.e., for both genotypes) for a given time-point then it was reported as absent and reported signal values and relative expression were ignored.

False Discovery Rate and p-values

The statistical method used to assign p-values to the fold-changes of gene responses is a two step procedure based on the Benjamini and Yekutieli construction of false discovery rate confidence intervals (FDRCI) (63-65) on the fold-changes between the $Nrl^{-/-}$ and wild type response profiles (55). FDRCI's are (1q) % confidence intervals where the level 'q' is corrected for error amplification inherent to performing multiple comparisons on many genes and many time points. For specified minimum fold-change (fcmin) and a given level of significance q, a gene response is declared as 'positive' if the range of the FDRCI is either greater than fcmin (positive fold-change) or less than -fcmin (negative fold-change). The FDRCI p-value for a given gene is defined as the minimum level q for which the gene's FDRCI does not intersect the interval [fcmin, fcmin]. For this data, we formed a ranked list of genes according to increasing FDRCI significance level having minimum fold-change (minfc) of 1.5 (0.58 log₂). All probesets with a p-value of less than 1 were reported.

Quantitative RT-PCR (Q-PCR)

RNA was treated with RQ1 DNAse (Promega, Madison, WI) following manufacturer's guidelines. Oligo-dT-primed reverse transcription was performed using 2.5 μ g of DNAse-treated total retinal RNA with Superscript II (Invitrogen, Carlsbad, CA). Primers for the validated genes were typically designed from the 3' UTR region using Primer 3 (http://www-genome.wi.mit.edu/cgi-bin/primer/primer3). PCR reactions on the cDNA template were then performed in triplicate in an I-cycler thermocycler with optical module (BioRad, Hercules, CA). Amplified products were quantified based on the level of fluorescence of SybrGreen I (Molecular Probes, Eugene, OR) in each reaction. Specificity of reactions was confirmed by melt curve analysis and gel electrophoresis. Average fold changes were then calculated based on the difference in the threshold cycles (C_t) between the NrI^{-/-} and wild type samples after normalization to *Hprt*.

Clustering Analysis

Clustering based on similarity of temporal expression profiles and visualization was performed using the software program Spotfire DecisionSite 7.2 (<u>www.spotfire.com</u>). The signal data of statistically significant differentially-expressed genes was standardized to z-scores (66), and hierarchical clustering performed using the "Euclidean distance" method.

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Annotation

Functional annotation of proteins was assigned through Gene Ontology (<u>http://www.geneontology.org</u>) or Locuslink (<u>http://www.ncbi.nlm.nih.gov/LocusLink</u>) classifications obtained through appropriate public databases such as NetAffx (<u>http://www.netaffx.com/indexp2.jsp</u>) (67) and DAVID http://apps1.niaid.nih.gov/david/upload.asp) (68).

Chromatin immunoprecipitation (ChIP) analysis

Retinas were obtained from C57BL/6 wild type mice and snap frozen on dry ice. ChIP was performed using a commercial assay kit (Upstate Biotechnologies, Charlottesville, VA). Briefly, four retinas were crosslinked in PBS containing proteinase inhibitors and a final concentration of 1% formaldehyde for 15 minutes at 37°C. The retinas were washed four times in ice-cold PBS with proteinase inhibitors and then incubated on ice for 15 minutes. The tissue was then sonicated on ice with ten 20 sec pulses. The remaining steps were performed as described by the manufacturer, using an anti-NRL polyclonal antibody (8).

The putative promoter region for each of the genes analyzed was determined using *in silico* methods (http://www.ncbi.nlm.nih.gov/mapview). Each promoter DNA sequence was analyzed using Matinspector (http://www.genomatix.de/index.html) and PCR primers were designed to flank putative AP1-like sites either predicted by Matinspector or predicted manually. If there was more than one AP-1 like site, the sequence element closest to the 5' untranslated sequence was used. Equal amounts of input DNA, with and without antibody, were used in each PCR reaction.

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Legends to Figures

Figure 1. Hierarchical clustering and functional annotation

Clustering of differentially expressed genes based on normalized average signals (z-scores). Bright green boxes indicate lowest signal with increasing values indicated by darkening color towards bright red, representing peak signal. The set of 161 non-redundant genes are divided, based on similarity of profiles, into 10 clusters. Down-regulated genes are in groups' I-VI (**A**), up-regulated in VII-X (**B**). Functional annotation is based on defined biological processes assigned by the Gene Ontology (GO) consortium. * Though listed as no defined function, Mtap6 is associated with microtubules and has a presumed role in synaptic plasticity and function. The far right column indicates whether the genes is known to be associated with (black square) or is a candidate for (empty square) retinal disease (see **Table 3** also).

Figure 2. Biological processes associated with differentially expressed genes

Upper panel shows overall distribution of all differentially expressed genes whereas lower panel a comparison between up and down-regulated genes.

Figure 3. ChIP analysis

PCR products are shown for 20 different gene promoters in which the most proximal putative AP-1/NRE-like site was assayed. Each set includes the input genomic DNA as positive control (lane 1), chromatin DNA fraction immunoprecipitated with anti-NRL antibody (lane 2), chromatin DNA obtained without the antibody (background control; lane 3) and water (negative control; lane 4). Enrichment (greater amount of product) with antibody IP over background control (no Ab) indicates the *in vivo* occupancy of a sequence element within the amplified region by Nrl. All assayed gene promoters showed enrichment, except Wisp1 (inconclusive) and Csda (negative). Direct regulation of Rho and Pde6b by Nrl (at these sites) has been previously demonstrated, and as such these are positive controls for the ChIP protocol.

Figure 4. Temporal expression of S-opsin in wild type and Nrl^{-/-} retina

The profile of relative expression of S-opsin determined by Q-PCR on the developing mouse retinas is shown after normalization to Hprt. Error bars indicate s.e.m.

Table 1a

Probeset	Acc#	Gene description	Name	P2 AFC	P10 AFC	2M AFC	P-value
96134_at	NM_139292	deleted in polyposis 1-like 1	Dp1I1	-	-10.12	-15.94	0.0001
96567_at	NM_145383	rhodopsin	Rho	-	-24.82	-41.30	0.0001
94701_at	NM_008806	phosphodiesterase 6B, cGMP, rod receptor, beta polypeptide	Pde6b	-2.29	-37.30	-25.00	0.0002
95389_at	XM_132106	cyclic nucleotide gated channel alpha 1	Cnga1	-	-7.25	-13.63	0.0004
93453_at	NM_009073	retinal outer segment membrane protein 1	Rom1	-2.17	-5.32	-3.27	0.0004
94853_at	NM_008142	guanine nucleotide binding protein, beta 1	Gnb1	1.17	-4.01	-8.12	0.0004
98531_g_at	NM_013525	growth arrest specific 5	Gas5	-1.15	-2.08	-6.28	0.0008
94139_at	NM_024458	phosducin	Pdc	-4.41	-3.84	-1.91	0.0008
93330_at	NM_007472	aquaporin 1	Aqp1	-	-1.33	-7.84	0.0011
104592_i_at	NM_025282	myocyte enhancer factor 2C	Mef2c	-	-	-4.18	0.0029
161871_f_at	NM_145383	rhodopsin	Rho**	-	-2.95	-7.80	0.0030
94150_at	NM_009118	retinal S-antigen	Sag	-5.07	-4.10	-1.20	0.0037
99860_at	NM_008140	guanine nucleotide binding protein, alpha transducing 1	Gnat1	-1.19	-55.61	-216.35	0.0046
95755_at	NM_011733	cold shock domain protein A	Csda	1.13	-1.51	-2.25	0.0059
161525_f_at	NM_009073	retinal outer segment membrane protein 1	Rom1**	1.04	-4.76	-3.70	0.0060
162167_f_at	NM_008806	phosphodiesterase 6B, cGMP, rod receptor, beta polypeptide	Pde6b**	-	-3.74	-4.99	0.0060
93533_at	NM_026899	RIKEN cDNA 1500011L16 gene	1500011L16Rik	-1.06	-2.62	-1.72	0.0065
93699 at		polymerase (DNA directed), gamma 2. accessory subunit	Pola2	-1.01	-1.73	-3.64	0.0069
93094 at	NM 007672	cerebellar degeneration-related 2	Cdr2	-1.07	-2.99	-5.47	0.0080
104591 g at	NM 025282	mvocvte enhancer factor 2C	Mef2c**	_	-	-2.53	0.0080
101923 at	NM 013737	phospholipase A2, group VII	Pla2o7	-1.03	-2.57	-1.82	0.0115
96831 at	NM 028295	protein disulfide isomerase-related	Pdir-pending	-1.02	-1.14	-2.68	0.0120
103895 at	NM 145930	expressed sequence AW549877	AW549877	1 03	-2.08	-1.30	0.0130
92796 at	NM 007431	alkaline phosphatase 2 liver	Akp2	-	-3 10	-3 17	0.0144
100696 at	NM 008805	phosphodiesterase 6A cGMP-specific rod alpha	Pde6a	_	-2.12	-2.95	0.0160
102890 at	NM 009228	syntrophin acidic 1	Snta1	-1 07	-2 74	-2.37	0.0194
160273 at	NM 007565	zinc finger protein 36. C3H type-like 2	Zfn36l2	1 41	-1.96	-3.68	0.0197
160134 at	XM 129394	adiponectin recentor 1	AdinoR1	-1.20	-1 42	-2 73	0.0248
98853 at	NM_008867	nhosnholinase A2 group IB nancreas, recentor	Pla2a1br	-	-3.13	-2.82	0.0240
100946 at	XM 286803	heat shock protein 1B	Hsna1h	_	_1 19	-3 17	0.0269
104590 at	NM 025282	myocyte enhancer factor 2C	Mef2c**	_	_	-3.66	0.0200
03875 at	XM 207062	heat shock protein 14	Hena1a	-1.05	_1 10	-2.62	0.0200
02601 at	NM 028302			-1.05	-3.10	-2.02	0.0230
92091_at	NM 007378	ATE binding cossotto sub family Λ (APC1) member 4		-1.41	-3.19	-1.20 0.00	0.0340
103733 of	AK013486			1.40	-2.70	1.06	0.0347
102044 of	NM 018865	WNT1 inducible signaling pathway protoin 1	2900000A00NIK	1.00	-1.43	-1.50	0.0303
102044_at	NM 008736		Nrl	-	2.50	-5.55	0.0479
102012_at	NM 000307	tumor pocrosis factor, alpha induced protein 3	Tofoio3	-	-2.55	-2.40	0.0510
99392_at	NM 120507	DNA comment Chr.6. Wound State University 176, every		-	-Z.1Z	-4.52	0.0560
90041_at	NM 052014	1 apulgiugeral 2 phoephote Q apultrangforage 2	Dowsul70e	1.00	-1.43	-3.27	0.0050
100007_at	NM 000795	r-acylgiyceror-s-priospriate O-acyltransierase 5	Agpais	-1.00	-1.10	-2.27	0.0700
90300_al	NM 009025	calcium channel, voltage dependent, alphaz/delta subunit 5	Cachazus	-	1.00	-2.00	0.0700
93390_g_at	NM 422825	prominin i		1.00	-1.80	-1.04	0.0810
1600097_at	NM 054007	DNA segment, Chr I, ERATO Doi 622, expressed	DIErtabZze	1.09	-1.33	-2.40	0.0934
160693_at	NM_054097	phosphatidylihositoi-4-phosphate 5-kinase, type II, gamma	Рірэк2с	-1.08	-1.90	-2.21	0.0980
9/090_at		serum/grucoconticola regulated kinase	Sgк	-1.09	-1.02	-2.69	0.0980
93328_at	NM_008230	nistiaine aecardoxylase	HOC	-	-	-2.12	0.1070
93887_at	NM_010820			1.06	-1.50	-2.60	0.1084
104206_at	NM_026153	RINEIN CUNA UD1UU1ZAU5 gene	0610012A05Rik	-2.07	-5.70	-2.66	0.1195
96596_at	NM_008681	N-myc downstream regulated-like	Ndri	-1.05	-1./8	-2.64	0.1370

102292_at	NM_007836	growth arrest and DNA-damage-inducible 45 alpha	Gadd45a	1.10	-2.15	-2.32	0.1450
160948_at	NM_008915	protein phosphatase 3, catalytic subunit, gamma isoform	Ррр3сс	1.01	-1.07	-2.27	0.1578
97770_s_at	NM_138587	DNA segment, Chr 6, Wayne State University 176, expressed	D6Wsu176e**	1.20	-1.27	-2.84	0.1610
160464_s_at	NM_010884	N-myc downstream regulated 1	Ndr1	-1.08	-1.82	-2.60	0.1819
94739_at	NM_011643	transient receptor potential cation channel, subfamily C, member 1	Trpc1	1.00	-1.13	-2.55	0.1948
97755_at	NM_007878	dopamine receptor 4	Drd4	-	-4.54	-2.46	0.2070
101151_at	NM_009038	recoverin	Rcvrn	-	-2.37	-2.12	0.2120
97357_at	AK044589	myocyte enhancer factor 2C	Mef2c**	-	-	-2.39	0.2140
96354_at	NM_020007	muscleblind-like (Drosophila)	Mbnl1	1.05	-2.02	-1.47	0.2530
95603_at	NM_138595	glycine decarboxylase	Gldc	-1.13	-2.15	1.17	0.2840
102413_at	NM_057173	LIM domain only 1	Lmo1	-1.16	-1.54	-1.87	0.2940
98993_at	NM_012023	protein phosphatase 2, regulatory subunit B (B56), gamma isoform	Ppp2r5c	-1.08	-1.45	-2.12	0.3280
93130_at	NM_183212	hypothetical protein D030064A17	D030064A17	1.04	-1.68	-1.96	0.3570
93389_at	NM_008935	prominin 1	Prom1**	1.01	-1.82	-1.64	0.3640
93202_at	NM_011851	5' nucleotidase, ecto	Nt5e	1.09	-2.07	-2.65	0.3720
95285_at	XM_128499	KRAB box containing zinc finger protein	KRIM-1	1.05	-1.12	-1.84	0.4030
102352_at	NM_134189	N-acetylgalactosaminyltransferase 9	Galnt9	1.03	-1.28	-1.80	0.4760
98569_at	NM_146118	mitochondrial Ca2+-dependent solute carrier	Mcsc-pending	-1.18	-1.47	-1.98	0.4830
99461_at	NM_008225	hematopoietic cell specific Lyn substrate 1	Hcls1	-	-	-1.90	0.4850
97579_f_at	NM_027010	crystallin, gamma F	Crygf	1.82	2.46	-3.51	0.4940
160808_at	NM_031869	protein kinase, AMP-activated, beta 1 non-catalytic subunit	Prkab1	-1.15	-1.55	-2.13	0.5120
101308_at	NM_008420	potassium voltage gated channel, Shab-related subfamily, member 1	Kcnb1	-	-2.14	-2.43	0.5120
103026_f_at	NM_007777	crystallin, gamma E	Cryge	1.74	2.24	-3.07	0.5220
98524_f_at	AK008780	RIKEN cDNA 2210039B01 gene	2210039B01Rik	1.02	-1.58	-2.26	0.5290
98329_at	XM_193953	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 2	Pfkfb2	-	-1.47	-1.77	0.5697
103460_at	NM_029083	HIF-1 responsive RTP801	Rtp801-pending	1.00	-1.17	-2.14	0.5980
100757_at	XM_194003	calcium channel, voltage-dependent, beta 2 subunit	Cacnb2	-	-2.23	-1.81	0.6290
102835_at	NM_007459	adaptor protein complex AP-2, alpha 2 subunit	Ap2a2	1.02	-1.52	-1.83	0.6490
99586_at	NM_009976	cystatin C	Cst3	-1.07	-1.71	-1.60	0.6940
96156_at	AK003573	RIKEN cDNA 1110008H02 gene	1110008H02Rik	-3.21	-2.38	-4.42	0.7627
104171_f_at	NM_007776	crystallin, gamma D	Crygd	1.83	2.37	-3.16	0.7652
96766_s_at	NM_019392	TYRO3 protein tyrosine kinase 3	Tyro3	-1.16	-1.07	-1.63	0.7750
101489_at	NM_009665	S-adenosylmethionine decarboxylase 1	Amd1	1.20	-1.04	-1.80	0.8460
103922_f_at	NM_028057	cytochrome b5 reductase 1 (B5R.1)	Cyb5r1-pending	1.11	-1.42	-2.04	0.8540
92790_at	NM_010655	karyopherin (importin) alpha 2	Kpna2	1.30	-1.35	-1.97	0.8670
92607_at	NM_008590	mesoderm specific transcript	Mest	1.41	1.08	-2.02	0.9058
101702_at	NM_011302	retinoschisis 1 homolog (human)	Rs1h	-	-2.70	-1.59	0.9617

Table 1b

Probeset	Acc#	Gene description	Name	P2 AFC	P10 AFC	2M AFC	P-value
92237_at	NM_009107	retinoid X receptor gamma	Rxrg	-1.01	2.50	2.92	0.0000
160893_at	NM_023121	guanine nucleotide binding protein, gamma transducing activity pp 2	Gngt2	1.14	6.20	7.90	0.0001
98807_at	NM_008141	guanine nucleotide binding protein, alpha transducing 2	Gnat2	1.48	7.20	7.58	0.0002
162287_r_at	NM_017474	chloride channel calcium activated 3	Clca3	-	3.20	14.88	0.0003
99395_at	NM_007538	opsin 1 (cone pigments), short-wave-sensitive	Opn1sw	1.29	8.43	9.40	0.0004
102151_at	NM_007419	adrenergic receptor, beta 1	Adrb1	1.16	4.33	4.55	0.0007
98498_at	NM_007611	caspase 7	Casp7	1.06	1.75	6.36	0.0017
98499_s_at	NM_007611	caspase 7	Casp7**	-1.05	1.67	5.93	0.0023
96920_at	NM_019564	protease, serine, 11 (Igf binding)	Prss11	1.13	2.30	1.36	0.0023
98427_s_at	NM_008689	nuclear factor of kappa light chain gene enhancer in B-cells 1, p105	Nfkb1	1.05	1.81	3.05	0.0037
103198_at	AI848576	expressed sequence Al848576	AI848576	-	2.37	1.52	0.0040
104346_at	NM_028250	acyl-Coenzyme A binding domain containing 6	Abdc6	1.06	1.70	2.67	0.0040
98918_at	NM_145367	thioredoxin domain containing 5	Txndc5	-1.09	1.17	2.69	0.0047
160754_at	NM_011224	muscle glycogen phosphorylase	Pygm	-	1.25	4.28	0.0050
98957_at	NM_023277	junction adhesion molecule 3	Jam3	1.03	2.03	3.31	0.0052
98967_at	NM_021272	fatty acid binding protein 7, brain	Fabp7	1.12	3.27	7.28	0.0065
95356_at	NM_009696	apolipoprotein E	Apoe	-1.16	2.23	1.70	0.0066
101855_at	NM_010837	microtubule-associated protein 6	Mtap6	-1.09	1.52	2.98	0.0070
104643_at	XM_109956	KIBRA protein	Kibra	1.13	5.51	5.98	0.0070
93482 at	AK020638	RIKEN cDNA 9530072E15 gene	9530072E15Rik	1.08	2.17	2.78	0.0080
	NM 007939	Eph receptor A8	Epha8	-1.04	2.65	1.22	0.0080
	 XM_148966	inhibin beta-B	Inhbb	1.26	2.83	-	0.0080
		quanylate cyclase activator 1a (retina)	Guca1a	-1.21	5.65	1.45	0.0115
99238 at	NM 013530	guanine nucleotide binding protein, beta 3	Gnb3	1.46	2.75	5.34	0.0130
98852 at	NM 178280	sal-like 3 (Drosophila)	Sall3	1.23	3.46	2.63	0.0130
93290 at	NM 013632	purine-nucleoside phosphorylase	Pnp	1.47	2.97	4.22	0.0139
100453 at	NM 007595	calcium/calmodulin-dependent protein kinase II. beta	Camk2b	1.01	2.30	3.11	0.0166
94338 g at	NM 008087	growth arrest specific 2	Gas2	1.19	10.07	1.78	0.0177
101344 at	NM 007627	cholecystokinin B receptor	Cckbr	1.04	1.44	2.34	0.0177
95363 at	NM 008504	granzyme M (lymphocyte met-ase 1)	Gzmm	-	3 16	4 82	0 0180
98560 at	NM 028870	clathrin light polypeptide (Lcb)	Cltb	1 0 1	2 43	3 71	0 0180
92904 at	NM 007548	PR domain containing 1 with 7NF domain	Prdm1	1 13	1 40	2 71	0.0300
102234 at	NM 024461	RIKEN cDNA 1810037117 gene	1810037I17Rik	1 89	1 78	2 62	0 0314
92293 at	NM 176930	neuronal cell adhesion molecule	Nrcam	1 16	1.62	2 27	0.0316
99111 at	NM 009191	suppressor of K+ transport defect 3	Skd3	1.03	2 25	2 23	0.0347
93973 at	NM 133916	eukarvotic translation initiation factor 3 subunit 9 (eta)	Fif3s9	-1.08	1 41	2 24	0.0370
97206 at	NM 016907	serine protease inhibitor. Kunitz type 1	Spint1		2 28		0.0380
98338 at	NM 010134	engrailed 2	En2	-1 02	1.04	2 72	0.0426
92415 at	NM 009404	tumor necrosis factor (ligand) superfamily, member 9	Tnfsf9	1.05	2.65		0.0479
101190 at	NM_008639	melatonin recentor 1A	Mtnr1a	1 18	2.33	_	0.0566
96911 at	XM 282613	guanine nucleotide binding protein, beta 2	Gnb2	-1.01	1.30	2 22	0.0566
160705 at	NM 007709	Chp/n300-interacting transactivator w/ E/D-rich carboxy-terminal dom 1	Cited1	1.01	2.34	1 55	0.0566
98424 at	NM 011204	notain tyrosine phosphatase, pon-recentor type 13	Pton13	1.00	1 54	2 24	0.0647
162206 f at	NM 007707	suppressor of cytokine signaling 3	Socs3	1 14	1.62	2.24	0.0650
06862 at	NM 13/05/		1110002B05Bik	1.14	2.28	2.44	0.0670
94256 at	NM 013885	chloride intracellular channel 4 (mitochondrial)	Clic4	1.27	1.65	2.45	0.0010
103033 at	NM 000780	complement component 4 (within H-2S)	C4	1.65	_1 11	2.20	0.0000
101706 at		evelic nucleatide gated channel alpha 3	Cnna3	-	2 2 /	2.03	0.0780
93269 at	NM 025374	alvoxalase 1	Glo1	1 00	<u> </u>	2.00	0 0000
50200_ut			0.01		1.771	2.10	0.0000

93731_at	NM_012056	FK506 binding protein 9	Fkbp9	-1.19	1.48	4.49	0.0910
96518_at	XM_109956	KIBRA protein	Kibra**	1.09	2.12	2.29	0.0934
103456_at	AW322500	expressed sequence AW322500	AW322500	-1.18	2.69	2.34	0.1190
92232_at	NM_007707	suppressor of cytokine signaling 3	Socs3**	1.06	1.64	2.37	0.1200
104374_at	NM_009252	serine (or cysteine) proteinase inhibitor, clade A, member 3N	Serpina3n	-	-	2.82	0.1251
104564_at	NM_009130	secretogranin III	Scg3	1.13	1.93	2.49	0.1251
103241_at	NM_153534	adenylate cyclase 2	Adcy2	1.34	1.92	1.95	0.1350
94393_r_at	NM_019423	ELOVL family member 2, elongation of long chain fatty acids (yeast)	Elovl2	1.33	2.82	3.09	0.1370
92534_at	NM_010276	GTP binding protein (gene overexpressed in skeletal muscle)	Gem	1.16	1.42	2.82	0.1535
100026_at	NM_007532	branched chain aminotransferase 1, cytosolic	Bcat1	1.01	1.38	1.96	0.1540
99972_at	NM_009414	tryptophan hydroxylase 1	Tph1	-	1.49	2.15	0.1590
95105_at	NM_025933	RIKEN cDNA 2010110M21 gene	2010110M21Rik	1.01	1.71	2.06	0.1670
97722_at	NM_025965	signal sequence receptor, alpha	Ssr1	2.17	2.03	1.96	0.1770
93268_at	NM_025374	glyoxalase 1	Glo1**	1.13	1.42	2.34	0.2070
93669_f_at	NM_009234	SRY-box containing gene 11	Sox11	3.41	1.75	-	0.2220
94872_at	NM_020561	acid sphingomyelinase-like phosphodiesterase 3a	Asml3a-pending	1.04	1.89	1.94	0.2220
99014_at	XM_133641	amyloid beta (A4) precursor protein-binding, family B, member 1	Apbb1	-1.01	1.58	1.88	0.2550
93412_at	NM_010045	Duffy blood group	Dfy	1.10	1.35	1.94	0.2570
97124_at	NM_008016	fibroblast growth factor inducible 15	Fin15	1.16	1.90	2.32	0.2940
104104_at	NM_030237	spermatogenic Zip 1	Spz1-pending	-1.20	1.20	2.41	0.3120
94733_at	NM_008830	ATP-binding cassette, sub-family B (MDR/TAP), member 4	Abcb4	1.05	1.37	1.77	0.3140
104761_at	XM_132228	anthrax toxin receptor	Antxr2	1.13	1.18	2.36	0.3275
101044_at	NM_008525	aminolevulinate, delta-, dehydratase	Alad	1.27	1.90	1.82	0.3328
99623_s_at	NM_019498	olfactomedin 1	Olfm1	1.05	1.47	2.18	0.3410
98544_at	NM_008193	guanylate kinase 1	Guk1	1.02	-1.01	1.88	0.3819
98111_at	NM_013559	heat shock protein 105	Hsp105	1.18	1.06	2.33	0.4600
92770_at	XM_192936	S100 calcium binding protein A6 (calcyclin)	S100a6	1.13	1.09	2.13	0.4870
101861_at	NM_011360	sarcoglycan, epsilon	Sgce	-1.03	1.38	1.70	0.4890
104388_at	NM_011338	chemokine (C-C motif) ligand 9	Ccl9	-	-	2.12	0.5800
160937_at	NM_016669	crystallin, mu	Crym	1.00	1.23	1.99	0.6290
93354_at	NM_007469	apolipoprotein C-I	Apoc1	-	1.82	-	0.6778
98005_at	NM_008862	protein kinase inhibitor, alpha	Pkia	1.07	1.25	2.14	0.6870
101191_at	NM_007873	double C2, beta	Doc2b	-1.01	-1.04	1.95	0.7286
104469_at	NM_010329	glycoprotein 38	Gp38	1.26	1.23	2.10	0.7560
94258_at	NM_007486	Rho, GDP dissociation inhibitor (GDI) beta	Arhgdib	1.09	1.08	1.86	0.7801
160414_at	NM_024249	RIKEN cDNA 1810073N04 gene	1810073N04Rik	1.01	1.84	1.36	0.8697
95397_at	XM_196081	RIKEN cDNA D430019H16 gene	D430019H16Rik	1.04	1.73	1.63	0.9320
93496_at	NM_134255	ELOVL family member 5, elongation of long chain fatty acids (yeast)	ElovI5	-1.02	1.26	1.96	0.9810

Table 1. Summary of microarray-based expression

Differentially expressed genes are listed based on statistical ranking (FDRCI p-values) and all are shown with a p-value <1 based on a 1.5 minimum fold-change. MGU74Av2 probeset IDs and corresponding Genbank accession numbers (RefSeq if available) are given. Where multiple probesets correspond to a single gene, the additional probesets are indicated by asterisks (**) and gray text. Average fold change (AFC) in expression in Nrl relative to wild type is shown for the three time-points analyzed, P2, P10 and 2-months (2M) after signal quantification and normalization by RMA. Gray dashes indicate that gene expression was reported as absent (by MAS5) on all eight GeneChips for the given time-point (i.e. wild type and Nrl^{-/-}). Gray numbers indicate a non-significant average fold change as determined by statistical analysis (p-value of 1). The lowest p-values are given for the three time-points measured for each of the gene probesets. Table **1a** shows down-regulated genes, **1b** up-regulated.

Table 2							
DOWN-I	REGULATE	D GENES		UP-RE	EGULATE	D GENES	
Name	AFC	qRT-PCR	P-value	Name	AFC	qRT-PCR	
Dp1l1	-15.9	-43.3	0.0001	Clca3	14.9	44.2	
Pde6b	-25.0	-52.7	0.0002	Opn1sw	9.4	23.1	
Rho	-41.3	-1604.0	0.0003	Casp7	6.4	10.7	
Cnga1	-13.6	-406.4	0.0004	Gnat2	7.6	15.3	
Gnb1	-8.1	-78.3	0.0004	Txndc5	2.7	7.8	
Gas5	-6.3	<u>-1.8</u>	0.0008	Fabp7	7.3	49.1	
Aqp1	-7.8	-52.3	0.0011	Epha8*	2.6	10.1	
Mef2c	-4.2	-4.6	0.0029	Inhbb*	2.8	9.5	
Gnat1	-216.4	-3494.4	0.0046	Rxrg	2.9	8.9	
Cdr2	-5.5	-7.0	0.0080	Guca1a*	5.6	5.5	
Pdir-pending	-2.7	-14.3	0.0120	Kibra	6.0	26.0	
Pde6a	-3.0	-13.3	0.0160	Gnb3	5.3	8.2	
Zfp36l2	-3.7	-7.7	0.0197	Sall3	2.6	2.4	
Rom1	-3.3	-5.7	0.0280	Camk2b	3.1	15.6	
2900026H06Rik*	-3.2	-4.7	0.0340	Gas2*	10.1	52.8	
Wisp1	-3.3	-32.7	0.0479	Sall3*	3.5	20.8	
Tnfaip3	-4.5	-7.5	0.0580	Gzmm	4.8	2.5	
D6Wsu176e	-3.3	-9.2	0.0650	Prdm1	2.7	7.6	
D1Ertd622e	-2.5	-7.7	0.0934	1810037I17Rik	2.6	2.1	
Abca4	-2.2	-6.6	0.1081	En2	2.7	31.3	
Mpdz	-2.6	-7.9	0.1084	Socs3	2.4	19.7	
0610012A05Rik	-2.7	-3.6	0.2140	Serpina3n	2.8	3.2	
Mbnl1*	-2.0	-2.2	0.2530	Elovl2	3.1	5.1	
Lmo1	-1.9	-9.6	0.2940	1110002B05Rik	2.4	<u>1.2</u>	
Drd4*	-4.5	-12.7	0.3410	Sox11**	3.4	<u>-2.0</u>	
Crygf	-3.5	-4.8	0.4940	Antxr2	2.4	7.0	
Crygd	-3.2	-6.5	0.7652	Olfm1	2.2	2.0	

Table 2. Real-time Q-PCR validation

Average Fold Change (AFC) is based on the microarray data with corresponding FDRCI p-value. Relative fold-change based on quantitative RT-PCR (Q-PCR) is shown. All measurements were on adult (2M) retina except where indicated (**P2, *P10). Underlined values indicate genes for which there are significant discrepancies between the two methods (i.e. no significant change reported by Q-PCR (<2-fold) or direction of change is in disagreement with microarray data).

KNOWN DISEASE GENES	Aver	age Fold	Change	Human	Homolog						
Mouse gene description	P2	P10	2M	Name	Location		DISEASE				
guanine nucleotide binding protein, alpha transducing 2	1.5	7.2	7.6	GNAT2	1p13.1		recessive achromatopsia (ACHM4)				
ATP-binding cassette, sub-family A (ABC1), member 4	-1.4	-2.8	-2.2	ABCA4	1p22.1-p21		recessive Stargardt (STGD1) / recessive MD / recessive RP (RP19) / recessive fundus flavimaculatus / recessive cone-rod dystrophy				
cyclic nucleotide gated channel alpha 3	-	2.2	2.6	CNGA3	2q11.2		recessive achromatopsia (ACHM2)				
retinal S-antigen	-5.1	-4.1	-1.2	SAG	2q37.1		recessive Oguchi disease / recessive RP				
guanine nucleotide binding protein, alpha transducing 1	-1.2	-55.6	-216.4	GNAT1	3p21		dominant congenital stationary night blindness				
rhodopsin	-	-24.8	-41.3	RHO	3q21-24		dominant RP / recessive RP / dominant congenital stationary night blindness				
cyclic nucleotide gated channel alpha 1	-	-7.3	-13.6	CNGA1	4p12-cen		recessive RP				
prominin 1	1.0	-1.9	-1.6	PROM1	4p15.33		recessive retinal degeneration				
phosphodiesterase 6B, cGMP, rod receptor, beta polypeptide	-2.3	-37.3	-25.0	PDE6B	4p16.3		recessive RP / dominant congenital stationary night blindness				
phosphodiesterase 6A, cGMP-specific, rod, alpha	-	-2.1	-3.0	PDE6A	5q31.2-q34		recessive RP				
guanylate cyclase activator 1a	-1.2	5.6	1.4	GUCA1A	6p21.1		dominant cone dystrophy				
opsin 1 (cone pigments), short-wave-sensitive	1.3	8.4	9.4	OPN1SW	7q31.3-q32		dominant tritanopia				
retinal outer segment membrane protein 1	-2.2	-5.3	-3.3	ROM1	11q13		dominant RP / digenic RP (with RDS)				
neural retina leucine zipper gene	-	-2.6	-2.5	NRL	14q11.1-q11.2		dominant RP (RP27)				
retinoschisis 1 homolog (human)	-	-2.7	-1.6	RS1	Xp22.2-p22.1		retinoschisis (XLRS1)				
CANDIDATE DISEASE GENES											
S100 calcium binding protein A6 (calcyclin)	1.1	1.1	2.1	S100A6	1q21	٦					
Duffy blood group	1.1	1.3	1.9	FY	1q21-q22	7	recessive cone-rod dystrophy (CORD8)				
retinoid X receptor gamma	-1.0	2.5	2.9	RXRG	1q22-q23	J					
adiponectin receptor 1	-1.2	-1.4	-2.7	ADIPOR1	1q32.1	ſ					
cytochrome b5 reductase 1 (B5R.1)	1.1	-1.4	-2.0	CYB5R1	1q32.1	7	recessive ataxia, posterior column with RP (AXPC1)				
guanylate kinase 1	1.0	-1.0	1.9	GUK1	1q32-q41	J					
nuclear factor of kappa light chain gene enhancer in B-cells 1, p105	1.1	1.8	3.1	NFKB1	4q24		Wolfram syndrome (WFS2)				
adenylate cyclase 2	1.3	1.9	2.0	ADCY2	5p15.3		dominant macular dystrophy (MCDR3)				
myocyte enhancer factor 2C	-	-	-4.2	MEF2C	5q14		dominant Wagner disease (WGN1) and erosive vitreoretinopathy (ERVR)				
5' nucleotidase, ecto	1.1	-2.1	-2.7	NT5E	6q14-q21		recessive RP (RP25)				
GTP binding protein (gene overexpressed in skeletal muscle)	1.2	1.4	2.8	GEM	8q13-q21	٦					
protein kinase inhibitor, alpha	1.1	1.2	2.1	PKIA	8q21.11	ſ	recessive optic atrophy (ROA1)				
LIM domain only 1	-1.2	-1.5	-1.9	LMO1	11p15	ך					
amyloid beta (A4) precursor protein-binding, family B, member 1	-1.0	1.6	1.9	APBB1	11p15	7	dominant atrophia areata; dominant chorioretinal degeneration, helicoid (AA)				
cholecvstokinin B receptor	1.0	1.4	2.3	CCKBR	11p15.4	J					
muscle givcogen phosphorylase	_	1.2	4.3	PYGM	11a13.1	ך					
microtubule-associated protein 6	-1.1	1.5	3.0	MAP6*	11a13.3	7	dominant neovascular inflammatory vitreoretinopathy (VRNI)				
suppressor of K+ transport defect 3	1.0	2.2	2.2	SKD3	11g13.3	J					
RIKEN cDNA 1110002B05 gene	1.2	2.3	2.4	C14orf147*	14g13 1	1	dominant MD. North Carolina-like with progressive sensorineural hearing loss				
purine-nucleoside phosphorylase	1.5	3.0	4.2	NP	14g13 1	ſ	(MCDR4) recessive rod monochromacy or achromatopsia (ACHM1)				
cerebellar degeneration-related 2	-1.1	-3.0	-5.5	CDR2	16p12.3		recessive RP (RP22)				
recoverin	-	-2.4	-2.1	RCV1	17p13 1	٦					
double C2. beta	-1.0	-1.0	2.0	DOC2B	17p13.3	7	dominant central areolar choroidal dystrophy (CACD)				
polymerase (DNA directed), gamma 2 accessory subunit	-1 0	-17	-3.6	POLG2	170	ſ					
guanine nucleotide binding protein, gamma transducing activity pp 2	1.1	6.2	7.9	GNGT2	17a21	7	cone rod dystrophy (CORD4)				
suppressor of cytokine signaling 3	1.1	1.6	24	SOCS3	17g25.3	J					
	1.1	1.0	-	20000	11920.0	-					

Table 3. Disease association or candidacy for differentially expressed genes

Differentially expressed genes that are known to be associated with retinal diseases or candidate genes that map to disease intervals are shown. Average fold change is shown as given in Table 1. *Indicates that gene is the likely human homolog but is not, as yet, definitively assigned as such (based on LocusLink).



1

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x

Sox11

x

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Figure 1





Figure 2



Figure 3



Figure 4

Abbreviations

Q-PCR = quantitative reverse-transcriptase coupled PCR RMA = Robust Multichip Average ChIP = chromatin immunoprecipitation GeneChip = Affymetrix GeneChip® AFC = Average Fold Change RP = Retinitis Pigmentosa FDRCI = False Discovery Rate Confidence Interval