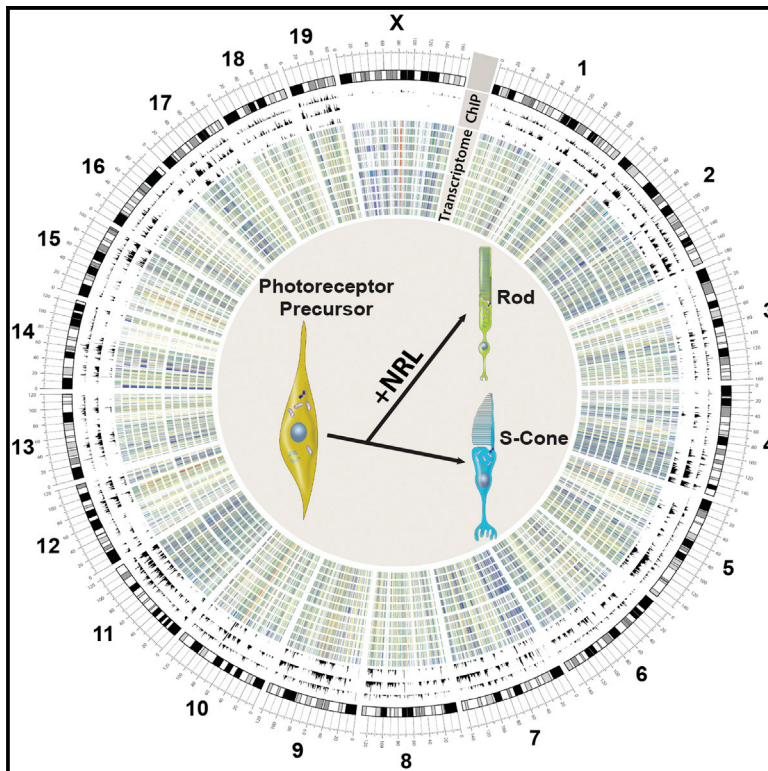


NRL-Regulated Transcriptome Dynamics of Developing Rod Photoreceptors

Graphical Abstract



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In Brief

A comprehensive integrated analysis of transcriptome and related NGS data by Kim et al. provides insights into rod photoreceptor morphogenesis and functional maturation, identifies previously unannotated transcripts, and reveals substructures of gene regulatory networks.

Highlights

- Rod photoreceptor transcriptome dynamics parallels rod morphogenesis and maturation
- Transcriptome transition during rod fate stabilization is disrupted by loss of NRL
- De novo assembly reveals previously unannotated rod-specific transcripts
- Integrated network analysis identifies secondary hubs of the NRL-centered rod GRN

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NRL-Regulated Transcriptome Dynamics of Developing Rod Photoreceptors

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SUMMARY

Gene regulatory networks (GRNs) guiding differentiation of cell types and cell assemblies in the nervous system are poorly understood because of inherent complexities and interdependence of signaling pathways. Here, we report transcriptome dynamics of differentiating rod photoreceptors in the mammalian retina. Given that the transcription factor NRL determines rod cell fate, we performed expression profiling of developing NRL-positive (rods) and NRL-negative (S-cone-like) mouse photoreceptors. We identified a large-scale, sharp transition in the transcriptome landscape between postnatal days 6 and 10 concordant with rod morphogenesis. Rod-specific temporal DNA methylation corroborated gene expression patterns. De novo assembly and alternative splicing analyses revealed previously unannotated rod-enriched transcripts and the role of NRL in transcript maturation. Furthermore, we defined the relationship of NRL with other transcriptional regulators and downstream cognate effectors. Our studies provide the framework for comprehensive system-level analysis of the GRN underlying the development of a single sensory neuron, the rod photoreceptor.

INTRODUCTION

Diversity in neuronal morphology and function is produced by coordinated and controlled changes in gene expression patterns

that are established by gene regulatory networks (GRNs), which provide roadmaps and topological information of regulatory and effector genes guiding specific biological processes (Peter and Davidson, 2011). Combinatorial actions of select transcriptional regulators on *cis*-regulatory genomic modules (Guillemot, 2007; Spitz and Furlong, 2012) generate context-, space-, and time-dependent gene expression states (Allan and Thor, 2015; Tsankov et al., 2015). The developmental transcriptome is dynamic and yields a deterministic framework for downstream physiological functions. Thus, comprehensive mapping of the transcriptomic landscape of a specific neuronal cell type can yield a better understanding of mechanisms underlying the acquisition and maintenance of its unique identity.

Retinal photoreceptors are sensory neurons that capture light and initiate the visual process (Lamb et al., 2007). Rod and cone photoreceptors have unique morphology to maximize their functional output, with numerous membrane discs in outer segments for photon capture and specialized ribbon synapses for signal transmission. Cones contain visual opsins with distinct absorption spectra and provide high-acuity daylight color vision, whereas rods mediate low-light vision using a single visual pigment, rhodopsin. Rod photoreceptors dominate the retina of most mammals (70%–80% of all retinal cells), including mice and humans, and are especially vulnerable to genetic changes in retinal and macular diseases (Wright et al., 2010).

Differentiation of rod photoreceptors spans a long temporal window far beyond their birth (~3 weeks in mice), thereby offering an attractive model to investigate transcriptome dynamics and GRNs that control their distinctive neuronal morphology and function. Several transcription factors (TFs), including OTX2, RORB, and PRDM1, regulate the photoreceptor lineage in developing retina (Brzezinski et al., 2013; Jia et al., 2009; Kato et al., 2010; Nishida et al., 2003; Roger et al., 2014); however, rod fate is critically dependent on expression and activity of

the Maf-family leucine zipper TF, NRL (Swaroop et al., 2010). Photoreceptor precursors fated to be rods acquire characteristics of short-wavelength cones (S-cones) in the absence of *Nrl* (*Nrl*^{-/-}) (Mears et al., 2001), whereas expression of *Nrl* in developing cones leads to rod differentiation (Oh et al., 2007). NRL collaborates with cone-rod homeobox CRX to activate rhodopsin and other rod-specific genes (Chen et al., 1997; Mitton et al., 2000). A key downstream target of NRL, the orphan nuclear receptor NR2E3, actively suppresses cone genes and consolidates the rod cell fate (Cheng et al., 2006; Oh et al., 2008). In addition, the estrogen-related receptor ESRRB, the myocyte enhancer factor MEF2C, the histone lysine-specific demethylase KDM5B, and the transcription-splicing protein NONO are among NRL targets that regulate specific aspects of rod development and survival (Hao et al., 2011, 2012; Onishi et al., 2010; Yadav et al., 2014). Thus, genome-wide targetome studies of NRL, CRX (Corbo et al., 2010; Hao et al., 2012), and other downstream TFs can be integrated with transcriptome profiles to assemble GRN modules associated with rod differentiation (Yang et al., 2015).

Here, we report directional RNA-sequencing (RNA-seq) analysis of developing mouse rod photoreceptors and *Nrl*^{-/-} photoreceptors fated to be rods, thereby directly determining the impact of perturbing a key hub in the rod-specific GRN. We tracked the dynamic changes occurring in the NRL-centered GRN and their impact on rod morphogenesis by integrating transcriptome, targetome, and DNA methylome datasets. We further define previously unannotated NRL-regulated transcripts by de novo assembly and assess the relevance of alternative splicing during photoreceptor development. Our studies thus provide a platform for system-level analysis of a single type of sensory neuron, the rod photoreceptor.

RESULTS

Transcriptome Profiling of Developing Rod Photoreceptors

In mice, the birth (final mitosis) of rod photoreceptors overlaps with the generation of all other retinal cell types and begins as early as embryonic day 12 (E12) (Figure 1A); however, a majority of rods are generated at postnatal day 0 (P0) to P2 (Akimoto et al., 2006; Carter-Dawson and LaVail, 1979). Although P2 mouse retina is enriched for newborn rods, robust upregulation of rhodopsin (encoded by *Rho*) and other phototransduction genes is detected later at P4. Outer segment formation and synaptogenesis begin at or after P6 (Blanks et al., 1974; Obata and Usukura, 1992) and continue through P10–P12. After eye opening at P14, synaptic changes are predicted based on activity until rods reach functional maturity by P21 (Figure 1A). To elucidate temporal dynamics of gene regulation, we generated rod photoreceptor transcriptome throughout development and then integrated differential expression analysis with NRL chromatin immunoprecipitation sequencing (ChIP-seq) data (Hao et al., 2012) and transcription factor binding motifs (Figure 1B).

Transcriptome profiling was performed using exon microarrays and directional RNA-seq platforms. We flow-sorted rods from *Nrlp*-GFP mouse retina (Akimoto et al., 2006) at P2, P4,

P6, P10, P14, and P28 for RNA-seq and P2, P4, P6, P8, P10, P12, and P21 for microarray analysis (Figures 1A, 1B, and S1). The two datasets demonstrated concordant transcriptome dynamics during differentiation. Principal-component analysis (PCA) revealed that developmental stage accounted for the largest variance among datasets in both platforms (Figure 1C). Biological replicates of each time point clustered together (Figure 1C), confirming high reproducibility. The rod transcriptome data were separated in an ordered fashion along PC1 and suggested a linear progression of gene expression changes during development (Figure 1C). The correlation between P21 microarray and P28 RNA-seq data (Figure 1D) indicated similar trends in the two platforms. Nonetheless, directional RNA-seq displayed a greater dynamic range, higher sensitivity in detecting expression changes (7,841 differentially expressed [DE] genes for P28/P2 RNA-seq having ≥ 2 -fold change, ≤ 0.01 false discovery rate [FDR], and ≥ 1 fragments per kilobase of transcript per million mapped reads [FPKM] at any time point vs. 2,533 DE genes with ≥ 2 -fold change and ≤ 0.05 FDR for P21/P2 microarray), and a negligible background noise compared to the hybridization-based microarray methodology. Therefore, we used directional RNA-seq data for subsequent comparative and integrated analysis. To facilitate rapid access, analysis, and visualization of datasets, we also developed an integrative platform called Retseq (<https://retseq.nei.nih.gov>).

Transition of Transcriptome Landscape in Developing Mouse Rods between P6 and P10

Hierarchical clustering and unsupervised Pearson's correlation analysis of averaged, normalized FPKM values at individual time points showed a decisive shift in rod transcriptome as a whole, and in specific transcripts with FPKM >5 in particular, between P6 and P10 (Figure 2A). A similar shift was apparent between P6 and P8 on PC1 analysis of microarray data (Figure 1C). Our data suggest that this transition in transcriptome landscape is highly synchronized and regulated, occurring within a relatively short time window, and constitutes a critical step in rod differentiation.

Specificity of the transcriptome data were validated by expression of known rod genes at P28 (Figure 2B). Furthermore, while a subset of cone genes were transiently expressed in rods at early time points, their expression was minimal or undetected by P28 (Figure 2B) (Kim et al., 2016). Pan-photoreceptor and pan-neuronal genes were highly expressed, whereas genes specific for inner retinal neurons (i.e., bipolar, ganglion, horizontal, and amacrine cells) and Müller glia were barely detectable in mature rods (Figure 2B).

Redundancy-removed Gene Ontology (GO) analysis (<https://david.ncifcrf.gov/>) of the top 1,000 highly expressed genes at each time point revealed both constitutive and stage-specific biological processes across development (data not shown). For example, early expressed genes were associated with RNA processing and metabolism, whereas photoreceptor development was among the highly enriched pathways at later stages. Enrichment of genes for ATP-synthesis-coupled proton transport and hexose catabolic process was consistent with high-energy requirement for photoreceptor morphogenesis and function (Kooragayala et al., 2015).

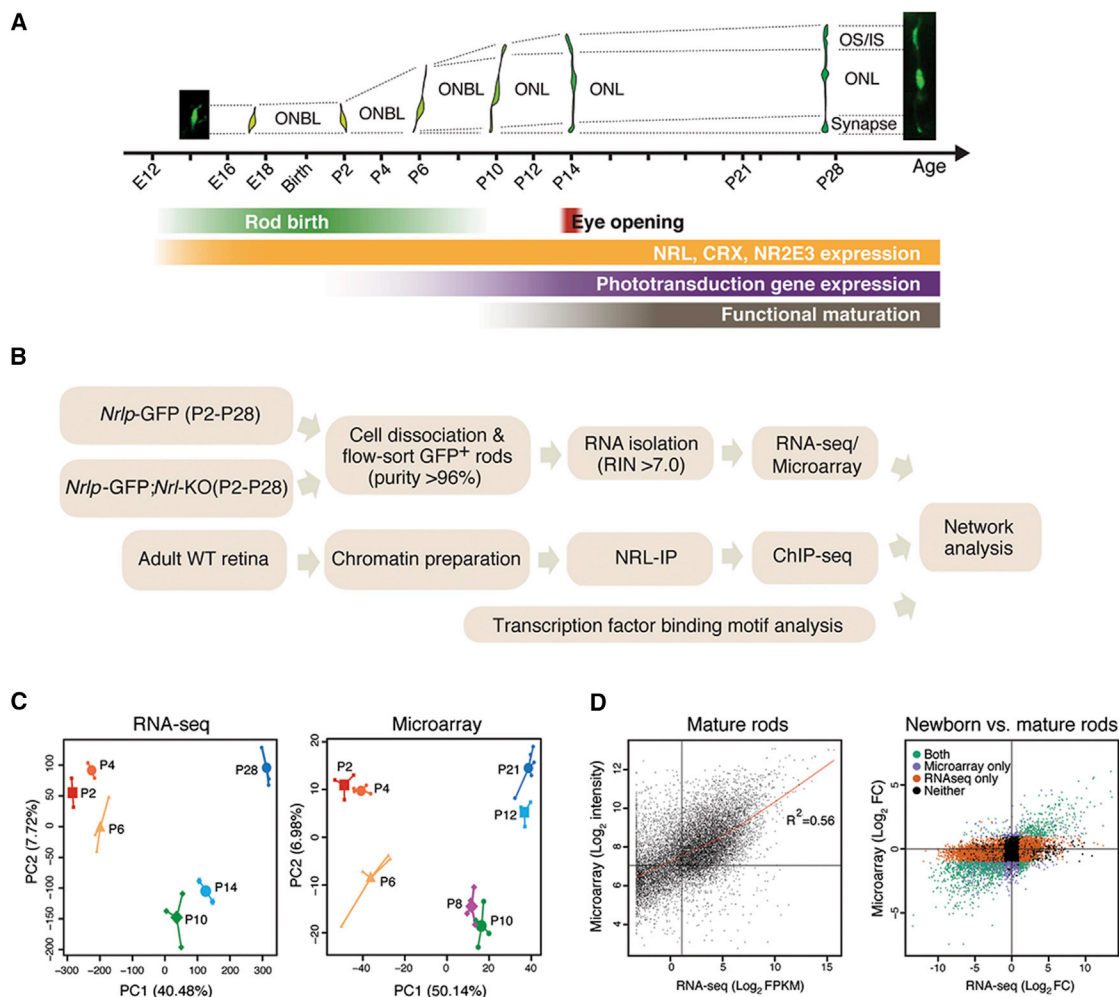


Figure 1. Study Design and Data Generation

(A) Time course of rod photoreceptor differentiation in mouse retina. Morphologies of developing and mature rods are shown at indicated developmental stages, with major events summarized below. ONBL, outer neuroblastic layer; ONL, outer nuclear layer; OS/IS, outer segment/inner segment.

(B) Flowchart of integrated transcriptome and network analysis. WT, wild-type; RIN, RNA integration number; IP, immunoprecipitation.

(C and D) Comparison of rod transcriptomes generated by directional RNA-seq and exon microarray. (C) Principal-component analysis (PCA) of directional RNA-seq (left) and microarray (right) data. The percentages indicate the amount of variation attributed to each principal component. Small shapes indicate individual samples, and larger shapes show the centroid of each grouping. (D) Correlation between RNA-seq and microarray data. RNA-seq and microarray data were de-duplicated and merged based on gene symbols. The highest FC representative from the RNA-seq transcriptome data were used as the gene representative for RNA-seq. Left: expression values (in \log_2 scale) of individual genes of mature rods (i.e., P28 in RNA-seq and P21 in microarray) were compared with each other using a scatterplot. \log_2 FPKM of one and \log_2 signal intensity of seven was used as a cutoff of actively transcribed genes in RNA-seq and microarray, respectively. The lowest regression line is shown in red, and the Pearson's correlation coefficient is indicated. Right: differentially expressed genes between newborn (P2) and mature rods (P21 for microarray and P28 for RNA-seq) were identified as having >2 -fold change and $FDR < 0.05$ in microarray data or < 0.01 in RNA-seq data. The scatterplot shows genes that are significantly differentially expressed genes in both analyses, microarray only, and RNA-seq only (green, purple, and orange dots, respectively). Black dots indicate genes with no significant change in expression in either analysis. FPKM, fragments per kilobase of exon model per millions of reads; FC, fold change; FDR, Benjamini-Hochberg false discovery rate.

Transcriptome of Developing S-Cone-like *Nrl*-KO Photoreceptors

To investigate the precise role of NRL in mediating transcriptome dynamics, we performed directional RNA-seq analyses of *Nrlp*-GFP;*Nrl*^{-/-} (or referred as *Nrl*-KO hereafter) photoreceptors (Akimoto et al., 2006). The data validated complete absence of rod-specific phototransduction genes and high expression of cone genes, as shown earlier by microarrays (Akimoto et al.,

2006; Yoshida et al., 2004). PCA analysis revealed a major divergence in transcriptome of S-cone-like *Nrl*-KO photoreceptors compared with wild-type rods (Figure 3A). The transcriptome of the two genotypes clustered at P2, but the difference increased dramatically as differentiation progressed (Figure 3A). Differential expression (DE) analysis identified 4,585 protein-coding transcripts (3,888 genes) showing significant up- or downregulation in *Nrl*-KO photoreceptors (FPKM > 5 in at least

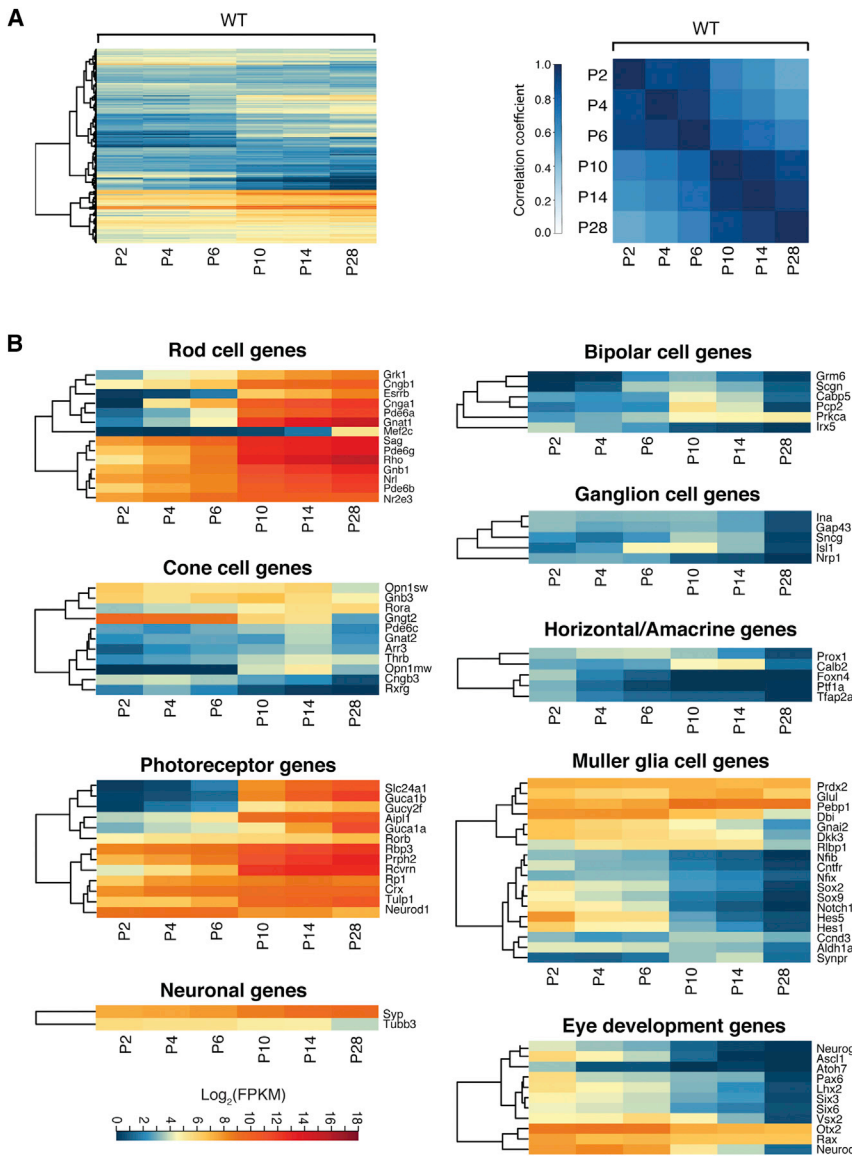


Figure 2. Transcriptome Dynamics during Rod Differentiation

(A) Hierarchically clustered heatmap of transcripts (a total of 11,146 transcripts that were expressed ≥ 5 FPKM at any time point in wild-type [WT]) during rod photoreceptor development (left) and correlation coefficient between every possible pair of transcriptomes at various differentiation stages calculated from directional RNA-seq data and presented in a matrix (right). A sharp change in transcriptome landscape is apparent between P6 and P10.

(B) Dynamic expression patterns of genes specific for photoreceptors (14 rod genes, 11 cone genes, and 13 genes expressed in both) and other retinal cell types during rod differentiation. In heatmaps, the average expression at each time point is plotted in \log_2 scale, and only those transcripts that were expressed at ≥ 5 in all replicates of at least one time point were included. Color scale is indicated in the bottom.

showed dramatically reduced expression in *Nrl*-KO photoreceptors (Figure 3D), suggesting its role in functional maintenance and/or survival of rods.

Stage-Specific DNA Methylation in Photoreceptor Genes

Rod-specific phototransduction genes, such as *Rho*, *Gnat1*, and *Cnga1*, exhibited a dramatic upregulation during P6–P10, and a subset of cone-specific genes were detectable in immature rods until a later stage (Figure 2B). To correlate DNA methylation to changes in global gene expression (Suzuki and Bird, 2008), we performed reduced representation bisulfite sequencing (RRBS) (Gu et al., 2011) using genomic DNA from P2, P10, and P28 rods. As predicted, the level of DNA methylation both in the gene promoter area and in the gene body showed a

negative correlation with gene expression at all three stages (Figure S2). A subset of rod-specific genes (e.g., *Rho* and *Pde6b*) showed high level of DNA methylation at their respective promoter at P2; however, by P28, the methylation was undetectable (Figure 4A). Interestingly, a greater number of rod genes had high level of DNA methylation in the gene body than in the promoter (Figure 4A). The degree of DNA methylation in the gene body of rod-specific genes was greatly reduced by P10–P28, consistent with delayed onset of their expression. As expected, *Nrl* exhibited constantly low DNA methylation at both promoter and gene body in rods at all times (Figure 4A). We further noted prominent DNA methylation in the gene body of cone-specific genes consistent with their lack of expression, while the promoter region of cone genes remained unmethylated (Figure 4A). Almost all non-photoreceptor retinal genes were unmethylated in their promoter region, though most genes acquired medium-to-high

one time point; fold change >2 ; FDR $<1\%$; Figure 3B). As anticipated from the PCA, the number of DE transcripts increased with rod morphogenesis and maturation (from 113 DE transcripts at P2 to 3,962 DE transcripts at P28). The dynamic transition in the transcriptome landscape from P6 to P10 that was apparent in rods was barely detectable in *Nrl*-KO photoreceptors (Figure 3B). To discriminate between direct and indirect targets of NRL, we integrated DE transcripts between rods and *Nrl*-KO photoreceptors with the NRL ChIP-seq data (Hao et al., 2012). As predicted, a subset of DE transcripts carried NRL ChIP-seq peaks (Figure 3C). Of these, 40 different genes encoded TFs that are putative direct targets of NRL (Figure 3D) and represent a starting point for potential secondary regulation functions. As an example, *Tnfrsf3*, previously implicated in negative regulation of cell death pathway, included two NRL binding sites and

negative correlation with gene expression at all three stages (Figure S2). A subset of rod-specific genes (e.g., *Rho* and *Pde6b*) showed high level of DNA methylation at their respective promoter at P2; however, by P28, the methylation was undetectable (Figure 4A). Interestingly, a greater number of rod genes had high level of DNA methylation in the gene body than in the promoter (Figure 4A). The degree of DNA methylation in the gene body of rod-specific genes was greatly reduced by P10–P28, consistent with delayed onset of their expression. As expected, *Nrl* exhibited constantly low DNA methylation at both promoter and gene body in rods at all times (Figure 4A). We further noted prominent DNA methylation in the gene body of cone-specific genes consistent with their lack of expression, while the promoter region of cone genes remained unmethylated (Figure 4A). Almost all non-photoreceptor retinal genes were unmethylated in their promoter region, though most genes acquired medium-to-high

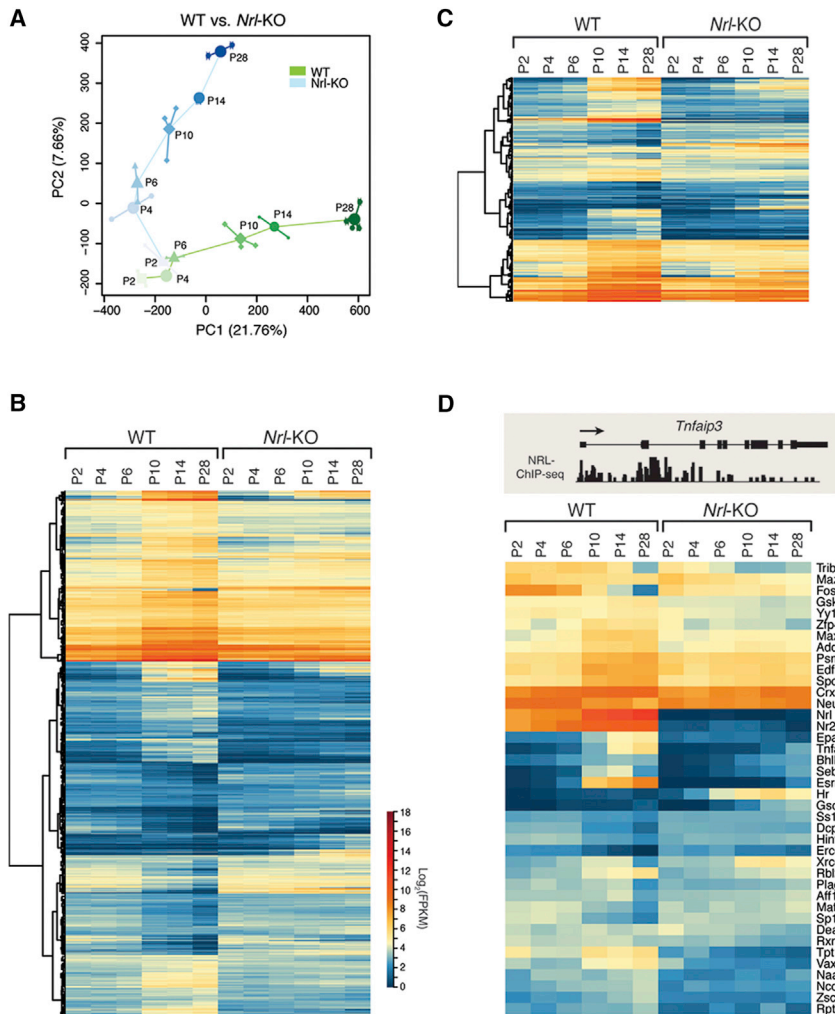


Figure 3. Transcriptome Regulation by NRL

(A) PCA analysis of time series RNA-seq data from *Nrl*-GFP+ wild-type rod photoreceptors and S-cone-like photoreceptors from the *Nrl*-KO retina. A significant change in transcriptome landscape was detected in GFP+ *Nrl*-KO photoreceptors.

(B) Heatmap of significantly, differentially expressed protein-coding transcripts. Color scale is indicated.

(C) Heatmap of differentially expressed genes containing NRL ChIP-seq peaks. An integrated analysis was performed combining differential gene expression data with the NRL ChIP-seq (Hao et al., 2012). The DE transcripts having at least one NRL ChIP-seq peak were considered putative, direct transcriptional targets and were included in the heatmap.

(D) Identification of transcription regulatory proteins that are putative NRL direct targets. Shown on the top is a genomic locus of one of the differentially expressed genes, *Tnfrsf3*, and NRL ChIP-seq coverage plot.

containing 775 putative conserved domains (E-val < 0.05). Notably, a sharp transition in expression between P6 and P10 rods was also evident in these transcripts (Figure 5A, left). However, the transition seemed more gradual, delayed or even absent in *Nrl*-KO photoreceptors (Figure 5A, right). We identified three major clusters based on expression patterns (Figure 5A). Members of cluster 1 showed a gradual increase in expression during rod differentiation, and this increase was not detected in the absence of NRL (Figure 5A), as is the case with rod-specific

DNA methylation in the gene body as rod maturation progressed (Figure 4B). No DNA methylation was detected in the promoter region of housekeeping genes, consistent with their constitutively high expression, with a few having medium level of DNA methylation in the gene body (Figure 4B).

De Novo Assembly and Identification of Previously Unannotated Transcripts

Our time series of developing mouse rods and S-cone-like *Nrl*-KO photoreceptors contained ~80,000 transcripts that were not annotated in the Ensembl database. Of these, over 20,700 constituted isoforms of known genes and 2,840 were multi-exonic intergenic transcripts (NIMETs) with an FPKM value ≥ 1 in all replicates of at least one time point (Figures 5A and S3). Mono-exonic transcripts and those expressed at <1 FPKM were excluded from further analysis, since these may represent transcriptional noise. Of 2,840 NIMETs, almost 15% showed <1 FPKM in RNA-seq data from other tissues examined (Consortium, 2004). Using four criteria (intergenic, multi-exonic, dynamic expression pattern, and at least one conserved domain), we identified 222 potentially protein-coding NIMETs (NPCs),

genes. A subset of transcripts in cluster 1 exhibited delayed onset of expression, suggesting their function in rod maturation. In contrast, transcripts in clusters 2 and 3 were highly expressed in *Nrl*-KO photoreceptors compared to rods (Figure 5A) and likely represented cone-enriched genes.

Conserved functional domains of NPCs were detected with HMMER against Pfam database (Finn et al., 2010). Identified domains spanned various functions from RNA processing (e.g., *Med19* and PF10278) to structural domains (e.g., *Golgin_A5* and PF09787) and protein degradation (e.g., *Peptidase_S49_N* and PF08496). Top represented domains (identified at least ten times) were grouped into families based on their putative function annotated in Pfam database. The most frequently occurring domains were of unknown function, but the second largest family was related to transcription regulation and included several zinc-finger domains (Figure 5B). Other highly represented families included PDZ and FERM domains involved in signaling and protein localization (Figure 5B). qRT-PCR analysis validated mature rod-specific expression of select NPCs (Figure 5C), arguing in favor of their functional relevance.

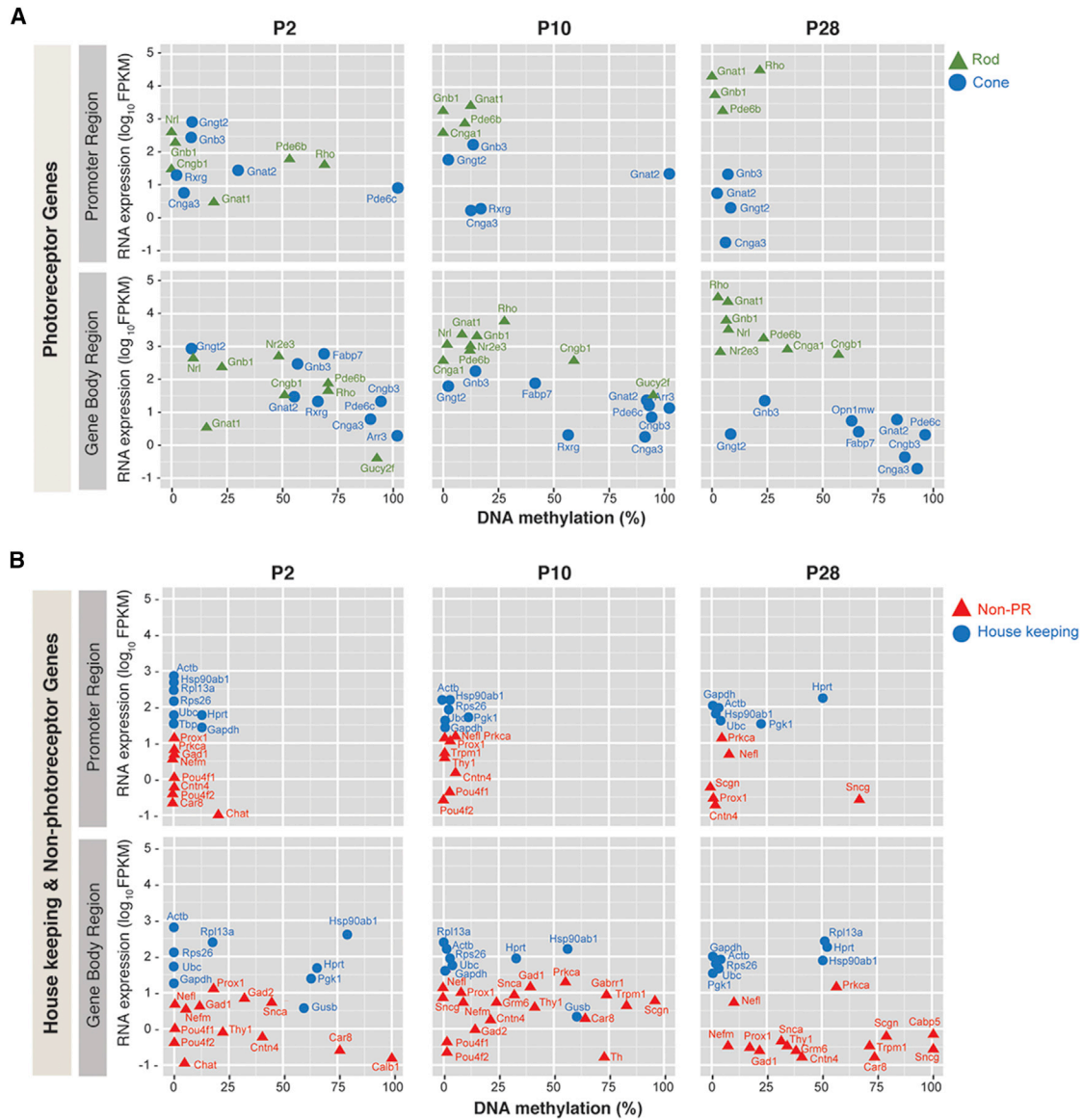


Figure 4. DNA Methylation of Photoreceptor Genes

(A) Correlation between DNA methylation level and gene expression of rod- and cone-specific genes. Degree of DNA methylation (percentage) in promoter or gene body region of individual rod- and cone-specific genes (green triangle and blue circle, respectively) are plotted against gene expression level (in \log_{10} FPKM). Please note that some promoters and gene bodies were not covered in RRBS analysis and thus not included in the plot.

(B) Plotted is correlation between DNA methylation level (percentage) in promoter or in gene body region and gene expression (in \log_{10} FPKM) of non-photoreceptor retinal genes (red triangle) and housekeeping genes (blue circle).

We superimposed genomic locations of NPCs with NRL and CRX ChIP-seq peaks using 1 kb as maximal allowed distance between the peak and the transcript start or end point. Of the 222 NPCs, 36 were potentially regulated by both NRL and CRX, while NRL alone regulated three and CRX 20 additional NPCs. Evolutionary conserved regions overlapped with predicted gene structure of 67% NPCs carrying both NRL and CRX ChIP-seq peaks (24 out of 36), with additional ten NPCs showing a partial overlap. For example, one NPC transcript, TCONS_00123129 (part of cluster 1), contained three significant Pfam domains (L27_1, PF09058; hDGE_aminylase,

PF14701; and CD36, PF01130) with strong evolutionary conservation in the gene structure, and its promoter region included NRL and CRX ChIP-seq peaks (Figure 5D).

As a proof of principle, we focused on TCONS_00068375, which is a part of potentially rod-enriched cluster 1 (Figures 5A and S3) and contains NRL and CRX ChIP-seq peaks. This transcript contains a G-patch domain (PF01585), present in RNA binding proteins, and is located between two annotated genes (*Slc22a28* and *Slc22a29*) that show very low or no expression in rods (Figure S4A). qRT-PCR and in situ hybridization (ISH) validated RNA-seq data and revealed its upregulation at P28

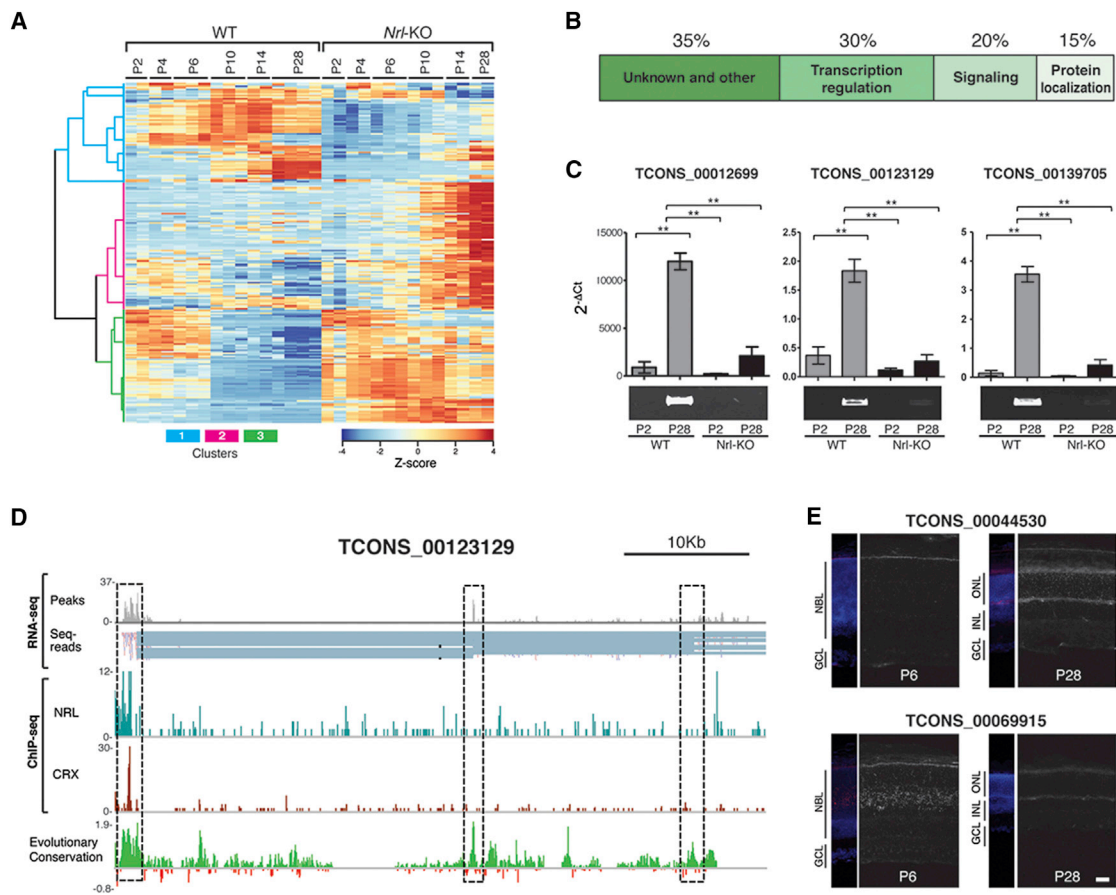


Figure 5. In Silico De Novo Analysis and Validation of Putative Protein Coding Transcripts

(A) Heatmap of standardized \log_2 FPKM values (with one offset) of 222 previously unannotated protein-coding transcripts (NPCs) across all time points in rods from wild-type and S-cone-like photoreceptors from *Nrl*-KO retina. Individual replicates are shown to display the degree of variability intrinsic to this type of analysis. The three largest clusters in the dendrogram are highlighted in blue (1), magenta (2), and green (3), respectively.

(B) Functional stratification of conserved protein domain identified by HMMER. Identified domains were grouped according to their described function in Pfam database. The relative frequency of occurrence was calculated for domains observed at least 10 times among the significant protein domain hits.

(C) qRT-PCR validation of three select NPC transcripts. Relative enrichment of each transcript compared with expression of housekeeping gene *Actb* was plotted for P2 and P28 wild-type and *Nrl*-KO retinas. Error bars indicate SEM. Double asterisks indicate statistically significant changes. Agarose gel images are shown below.

(D) Schematic representation of a NPC, TCONS_00123129. NRL and CRX ChIP-seq peaks are located in close proximity to the transcription start site (TSS) of this transcript, and placental mammalian evolutionary conservation peaks (as available from USCS genome browser) overlap with the predicted gene structure, highlighted by a dashed box. Histograms for RNA-seq and NRL and CRX ChIP-seq indicate the coverage of all aligned reads across the genomic area. The individual sequence reads are shown below the RNA-seq histogram. Red or blue blocks indicate discrete reads in antisense or sense orientation, respectively, with thin blue lines marking the splice portions of the reads.

(E) In situ hybridization of the previously unannotated transcripts TCONS_00044530 and TCONS_00069915 at P6 and P28 in wild-type retina. The signal for both transcripts follows the dynamic expression pattern seen in (A) and Figure S3. Blue, DAPI nuclear stain; red, in situ signal in merged images; white, in situ signal in single channel images. Scale bar, 50 μ m. GCL, ganglion cell layer; INL, inner nuclear layer; NBL, neuroblastic layer; ONL, outer nuclear layer.

(Figures S4B and S4C); this increase was not observed in *Nrl*-KO retina (Figure S4B). In vivo knockdown of TCONS_00068375 transcript in developing mouse retina resulted in thinning of outer nuclear layer (ONL) (Figure S4D), suggesting its potential function in rod photoreceptors.

ISH of two additional NPCs recapitulated their dynamic expression pattern in the ONL (Figure 5E). Although TCONS_00044530 was undetectable at P6 but observed at P28, TCONS_00069915 transcripts were prominent at P6 but absent at P28 (Figure 5E). TCONS_00044530 transcript also belongs to

cluster 1 (Figures 5A and S3) and is predicted to encode a protein with a nuclear RNA-splicing-associated domain (SR-25, PF10500), whereas TCONS_00069915 (cluster 3) likely encodes endonuclease/exonuclease/phosphatase family (Exo_endo_phos, PF03372) and syntaxin 6 (PF09177) protein domains (Figures 5A and S3).

Alternative Splicing and Alternate Promoter Usage

By surveying developmental time- and NRL-dependent alternative splicing (AS) events in RNA-seq data, we identified 809 and

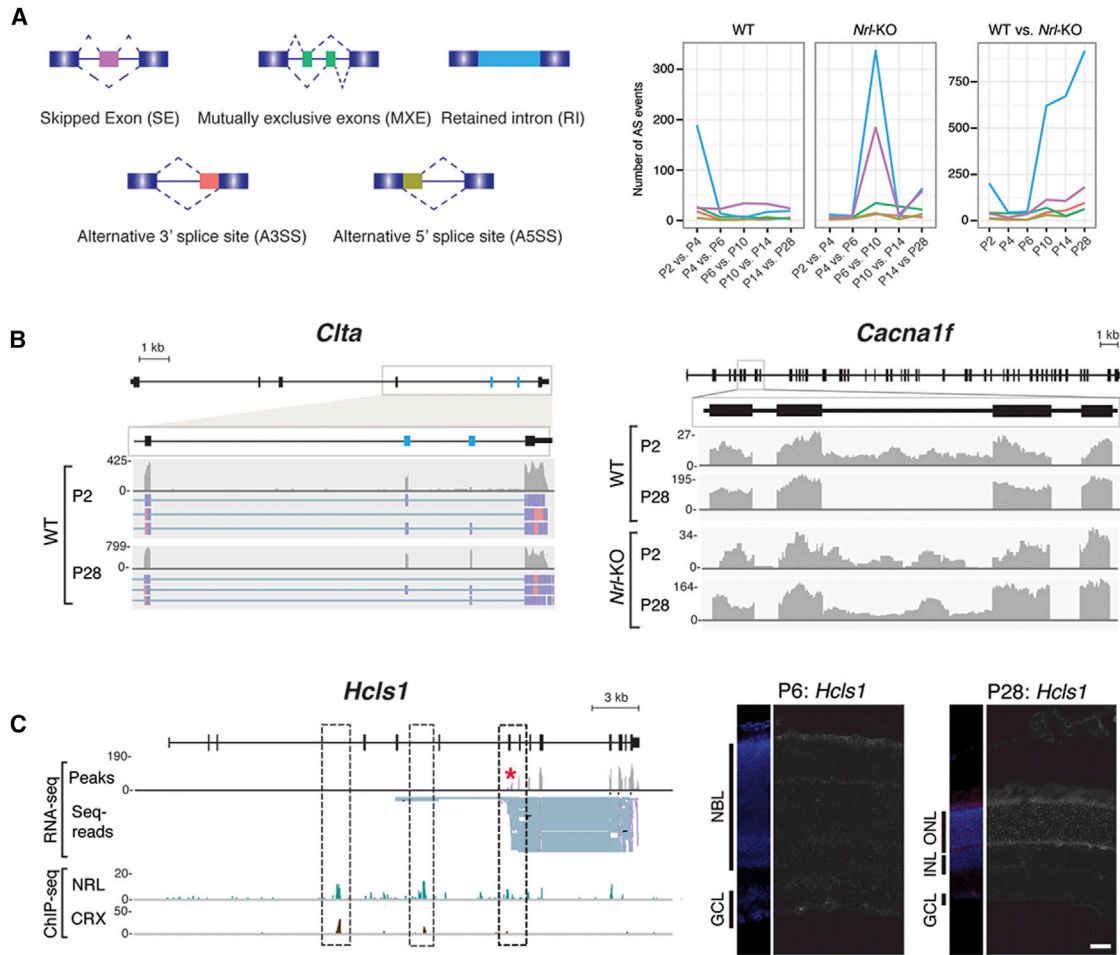


Figure 6. Alternative Splicing and Promoter Usage Events Identified during Rod Differentiation

(A) Types of alternative splicing events detected and their frequencies in time-wise and group-wise comparisons. Splicing events are color coded to match the relevant graphs.

(B) Differential splicing events during maturation of rods and *Nrl*-KO S-cone like photoreceptors. Shown is the genomic area of *Clta* gene (upper lane in the left panel), which comprises seven exons and six introns, and RNA-seq coverage plots of P2 and P28 wild-type rods in the area of exons four through seven (bottom lanes in the left panel). Solid boxes and lines represent exons and introns, respectively, and untranslated regions are illustrated as thinner boxes. The fifth and sixth exons of *Clta* gene are differentially spliced and highlighted in blue. Individual spliced reads demonstrate the splice isoform that are generated from *Clta* at both time points. Translated exons, untranslated exonic regions and introns are indicated as thick boxes, thin boxes and lines, respectively. RNA-seq coverage plots are shown as gray histograms. For *Clta* gene, individual sequence reads (red and blue boxes for antisense and sense orientation) with spliced area (thin blue lines) are also indicated.

(C) Alternative promoter usage of *Hcls1*. RNA-seq reads show only partial coverage of known exons of *Hcls1* (exons 8–14), with another exon introduced upstream to exon 8 (red asterisk). Dashed boxes indicate NRL and CRX ChIP-seq peaks. In situ hybridization for a retinal isoform of *Hcls1* (probe designed to hybridize to approximately 1000 bases starting in the previously un-annotated exon) shows significant increase in expression from P6 to P28. DAPI nuclear stain is indicated in blue, and in situ signal is indicated in red and in white in merged or in single channel images, respectively. Scale bar, 50 μ m. GCL, ganglion cell layer; INL, inner nuclear layer; NBL, neuroblastic layer; ONL, outer nuclear layer.

483 differential splicing events during the development of rods and *Nrl*-KO photoreceptors, respectively (FDR < 0.01; Figure 6). A comparison of all temporal data from rods versus *Nrl*-KO photoreceptors identified 3,644 differential AS events of all five major types (FDR < 0.01; Figure 6A) as well as potential alternative promoter usage and introduction of new exons. Rods and *Nrl*-KO photoreceptors revealed distinct genome-wide AS events. While transcripts with a retained intron (RI) were frequently detected in rods at P2 and P4, *Nrl*-KO photoreceptors showed increased RI and skipped exon (SE) between P6 and P10 (Figure 6A). The

most dramatic shift in AS was evident in rods versus *Nrl*-KO photoreceptors from P6 to P10 (Figure 6A), similar to that detected in the broader transcriptome landscape (see Figure 2A). RI was the most common AS event in rods with highest number at P28 (Figure 6A).

The two most frequent AS events observed, SE and RI, are exemplified by two genes, clathrin light polypeptide (*Clta*) and calcium channel, voltage-dependent, L type, α 1F subunit (*Cacna1f*), respectively (Figure 6B). The *Clta* gene produced three isoforms by skipping or including exons 5 and 6. In P2

rods, the two most abundant isoforms skipped either both or only exon 6; however, increased expression of exon-6-containing transcripts was observed at P28 (Figure 6B, left). The *Cacna1f* transcript in mature rods at P28 did not include any intron 6 sequences, which were evident at P2; however, introns were not removed in *Nrl*-KO photoreceptors, suggesting a role of NRL in transcript maturation (Figure 6B, right).

Consistent with previous observations (Hao et al., 2011, 2014), we identified several rod-specific alternative transcript isoforms that are regulated by NRL. Hematopoietic cell-specific Lyn substrate 1 (*Hcls1*), associated with gene regulation in the hematopoietic system (Skokowa et al., 2012), exhibited rod-specific, late-onset expression of a transcript which is initiated upstream of exon 8 from a new transcription start site, suggesting alternative promoter usage (Figure 6C). Furthermore, *Hcls1* included NRL and CRX ChIP-seq peaks upstream of a retina-specific exon. ISH demonstrated *Hcls1* expression specifically in the retinal ONL at P28 (Figure 6C), implying an as-yet-unrecognized function of this transcription factor in rod photoreceptors.

Co-expression Network Analysis

A network of transcription factors including NRL and CRX is predicted to drive the gradual transcriptome shift during rod maturation. By employing weighted gene co-expression network analysis (WGCNA) (Langfelder and Horvath, 2008) to all transcripts expressed at ≥ 5 FPKM at one or more time points, we identified 11 gene modules that correlated to distinct temporal expression patterns (Figure S5). A vast majority (672 out of 803) of TF-encoding transcripts was present in modules A, B, and C, which included all known rod-specific transcriptional regulators (Figure S5). Analysis of module A yielded a subcluster of 12 TFs including *Esrrb* and *Mef2C* (Figure S5, rod cluster), which are known regulators of rod differentiation (Hao et al., 2011, 2012; Onishi et al., 2010). The “rod cluster” exhibited a clear shift in expression between P6 and P10, with no or low expression in the early postnatal days and a substantially higher expression by P10 and later. This shift was not detected in *Nrl*-KO data.

NRL-Centered Transcriptional Network

To construct the GRN associated with rod morphogenesis and functional maturation, we integrated differential gene expression analyses of time series and knockout samples to NRL targetome data and transcription factor binding motifs. We noted that a subset of early NRL targets were transcriptional regulators (see Figure 3D), which in turn could modulate expression of other downstream genes. We thus investigated potential secondary hubs (referred as source genes herein) that cooperate with and mediate distinct subsets of NRL function. We identified four potential secondary hubs of the NRL-centered GRN—BHLHE41, ESRRB, FOS, and NR2E3 (Figure 7A)—by differential expression analysis comparing rods and *Nrl*-KO photoreceptors during the P2–P6 period and using a stringent cutoff of statistical significance and expression level (see Supplemental Experimental Procedures for details). The four secondary nodes exhibited distinct expression profiles during rod differentiation and upon genetic loss of *Nrl* (Figure 7B). To construct the NRL-centered network with the four secondary hubs, potential downstream tar-

gets of source nodes were identified using the JASPAR database of DNA-binding motifs (Sandelin et al., 2004). We confined the targets of source genes to those differentially expressed in P10, P14, and P28 rods compared to P6 or earlier. The reconstructed network had 1,015 nodes (i.e., individual genes that constitute the network) and 1,414 edges (i.e., predicted interaction among any two nodes within the network) (Figure 7C, right). Notably, phototransduction, visual perception, eye morphogenesis, photoreceptor cell development, and non-motile primary cilium (photoreceptor outer segment) were highly represented GO terms among the nodes in NRL-centered network (Figure 7C).

DISCUSSION

Neuronal cell-type-specific analysis has been limited to heterogeneous transcriptome landscapes of discrete regions of the brain (Hawrylycz et al., 2012; Tebbenkamp et al., 2014) or stem cell-derived neurons (Kaewkhaw et al., 2015; Thakurela et al., 2015; van de Leemput et al., 2014). Only a few neuron-specific transcriptomes have been reported so far (Akimoto et al., 2006; Macosko et al., 2015; Siegert et al., 2012; Telley et al., 2016; Zhang et al., 2014). Here, we used the mouse rod photoreceptor as a model sensory neuron to establish temporal transcriptome dynamics driven by the rod cell fate determination factor NRL, using two distinct platforms (RNA-seq and exon arrays) for cross-validation and deeper coverage of differentially expressed genes. We integrated multiple genome-wide expression datasets, before and after perturbing *Nrl*, and superimposed these with *cis*-regulatory maps obtained by NRL ChIP-seq (Hao et al., 2012), motif analysis and global DNA methylation profile. A genome level topology of “rod regulome” (Figure S6) provides a comprehensive view of the power of system-level analysis and a framework upon which additional “omics” and epigenetic data can be superimposed.

In this study, we focused on three fundamental questions: (1) are there strategic attributes in the transcriptional dynamics of developing rods; (2) do we detect unknown or unusual elements (genes or transcripts) that complement the existing rod transcriptome landscape; and (3) how does the NRL-centered network expand to complete rod development? The most notable characteristic we identified is the dramatic shift in transcriptome profile between P6 and P10 in developing rods. Postmitotic photoreceptors appear to retain developmental plasticity (Adler and Hatlee, 1989; Brzezinski et al., 2013; Cheng et al., 2006; Ng et al., 2011; Oh et al., 2007) until their fate is stabilized by additional changes in the differentiation program. The observed change in rod transcriptome between P6 and P10 may represent the decisive transition from early developing rods (still plastic) to functionally maturing rods (with stable cell fate). Downregulation of *Vsx2* after P10 (see Figure 2B) is consistent with its proposed role in controlling bipolar cell genesis through inhibition of rod fate (Livne-Bar et al., 2006). Similarly, PRDM1, which regulates *Otx2* and the decision between rod and bipolar fate in some progenitors (Wang et al., 2014), is also downregulated by P10. Low-level expression of *Vsx2* and *Prdm1* in early developing rods may indicate a molecular memory from the original retinal progenitor. Notably, we detected

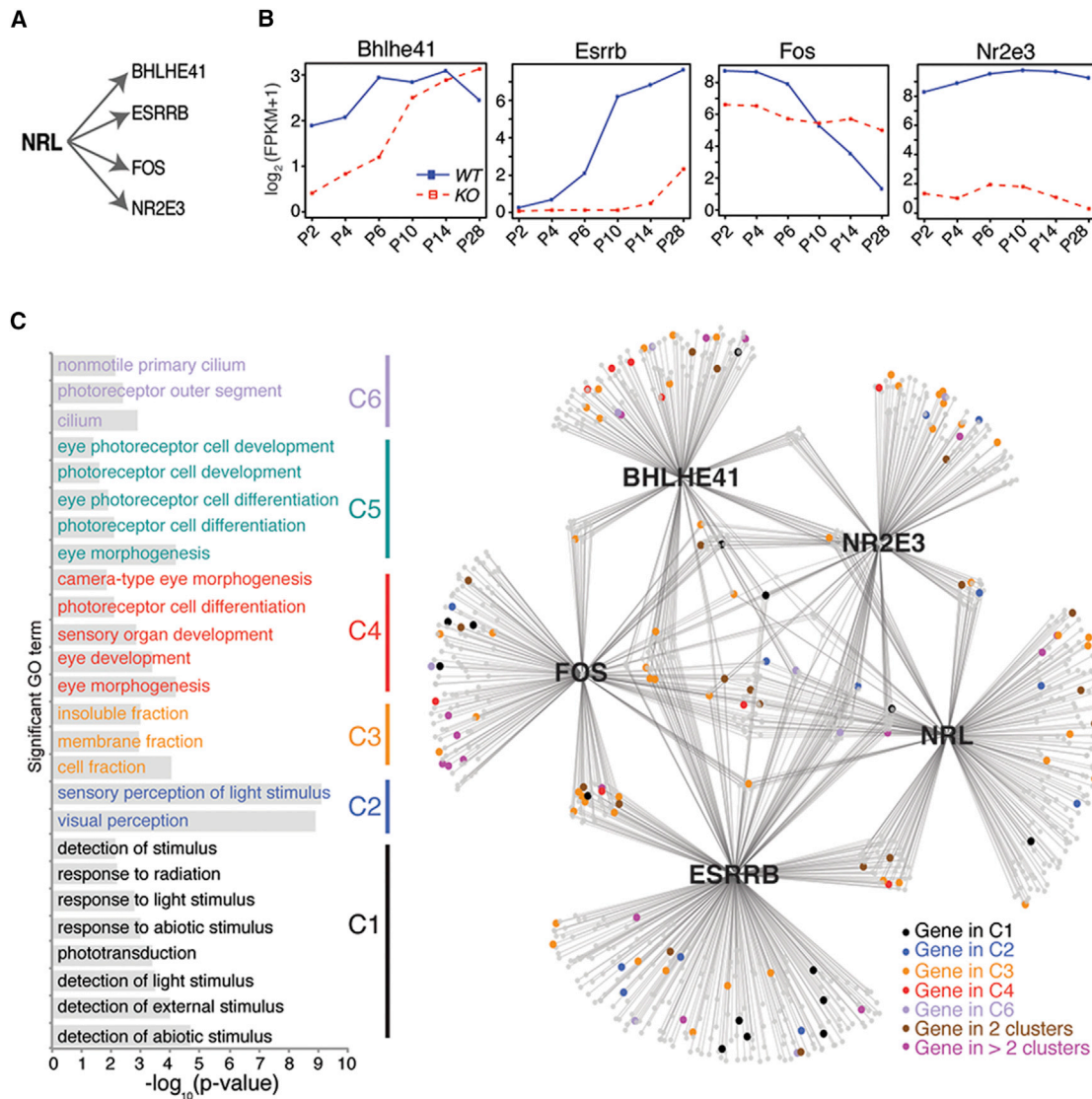


Figure 7. Gene Regulatory Network Analysis

(A) Regulatory hubs of NRL-centered GRN.

(B) Temporal expression profile of secondary hubs. Expression values of four genes in rods (blue line) and *Nrl*-KO photoreceptors (red dashed line) are plotted in log₂ scale for each time point. When multiple transcripts are expressed, expression level of the most highly expressed transcript is indicated.

(C) NRL-centered network and highly represented GO terms among target genes of NRL and secondary hubs. Circles indicate individual nodes (i.e., regulators and target genes) with primary regulatory hub NRL and four other secondary hubs labeled, and lines indicate predicted interaction between regulatory hubs and targets. Some targets are regulated by one regulatory gene, while others have regulatory interaction by two or more hubs. Target genes that belong to highly represented GO terms grouped into C1–C5 were highlighted in the indicated color.

expression of a number of S-cone genes in developing but not in mature rods (Kim et al., 2016), consistent with the hypothesis of S-cone as a “default” fate in many (likely prenatal) photoreceptor precursors in mouse retina (Swaroop et al., 2010) and suggesting the recruitment of S-cones to rod fate to mitigate nocturnal bottleneck during early mammalian evolution (Kim et al., 2016). We believe that the P6 to P10 transition in rod transcriptome consolidates rod versus bipolar or rod versus cone alternative fates by decisively suppressing molecular footprints associated with early steps in lineage selection.

NRL is the primary regulatory node in the rod GRN and orchestrates the consolidation of rod fate, expression of phototransduction genes, outer segment morphogenesis, assembly of pre-synapse, and functional maturation by directly or indirectly modulating the expression of most photoreceptor-enriched genes (Hao et al., 2012). Our analysis reveals previously unappreciated substructures of the NRL-centered network with four putative secondary hubs: BHLHE41 (identified by the WGCNA analysis as well), ESRRB, FOS, and NR2E3. Identification of NR2E3 provides credibility to other predicted secondary hubs.

Interestingly, 1,015 genes that comprise the new network segregate into distinct clusters that are enriched in genes associated with membrane, cilia and outer segments, as well as phototransduction. These genes are differentially expressed at or after P10 and are either direct targets of NRL or are indirectly regulated through the secondary hubs. Unlike the consistently high expression of NRL, three of the secondary hubs (BHLHE41, ESRRB, and FOS) exhibit stage-dependent expression patterns, which may allow for a stepwise regulation and explain delayed onset of rod phototransduction gene expression and cilia/synapse formation. Of note, BHLHE41 is shown to participate in regulating circadian rhythm-associated genes, like *PER1*, which in turn may impact circadian-associated rod physiology and function (Green and Besharse, 2004; McMahon et al., 2014).

Control of rod development by NRL is likely fine-tuned by tertiary and higher-order co-regulators and chromatin accessibility (Spitz and Furlong, 2012). Co-expression and network analyses reveal additional regulatory proteins that show a sharp increase in expression between P6 and P10, suggesting their possible role as co-regulators with NRL in controlling subsets of targets to produce rod-specific attributes. Known NRL-interacting TFs (MEF2C and ESRRB) were identified by the WGCNA analysis, together with potential regulators (HCLS1 and BHLHE41), expanding the repertoire of regulatory machinery involved in photoreceptor development.

In our study, the probability to detect true positives by genome-guided de novo assembly (Li et al., 2014) was augmented by temporal rod transcriptome analysis together with the expression data from *Nrl*^{-/-} S-cone-like photoreceptors. Many previously un-annotated transcripts are predicted to encode proteins with DNA-binding zinc-finger Cys2-His2 (zf-C2H2) domains that might have significant regulatory functions (Najafabadi et al., 2015) in fine-tuning photoreceptor maturation. ISH and qRT-PCR validation of select transcripts and the presence of NRL and CRX ChIP-seq peaks (see Figures 5C–5E) seem to corroborate our hypothesis. Indeed, knockdown of TCONS_00068375 showed thinning of the outer nuclear layer, consistent with its expression pattern and regulation by NRL. Our analysis also validated the prevalence of AS in developing photoreceptors, as implicated by whole-retina studies (Liu and Zack, 2013; Wan et al., 2011). We identified a number of previously unannotated transcript isoforms and recognized RI as the dominant AS event in developing photoreceptors. Our study corroborates the reported importance of RI in differentiating mammalian cells (Wong et al., 2013; Yap et al., 2012) and further uncovers a potential role of NRL in facilitating appropriate splicing of gene products in maturing rods.

The transcriptome catalog of rod development and associated gene regulatory network provides a reference map to investigate photoreceptor biology with potential implications for human retinal degeneration. Despite the identification of over 200 genes associated with photoreceptor disease (RetNet; <https://sph.uth.edu/retnet/>), not enough is known about the precise impact of disease-causing mutations on gene/protein networks controlling photoreceptor homeostasis. The existence of non-coding genetic variants in the human population (1000 Genomes Project Consortium et al., 2015) and those specifically segregating in Mendelian diseases (Cartegni et al., 2002) poses a challenge in

establishing genotype-phenotype relationships. The problem is even more acute in complex multifactorial diseases (e.g., age-related macular degeneration) where a vast majority of associated variants are in the non-coding region (Chakravarti et al., 2013; Fritsche et al., 2016). Previously unannotated transcript isoforms and/or intergenic transcripts identified here may provide insights in the missing heritability.

Many complex neurodegenerative diseases afflict distinct subsets of neurons (Mattson and Magnus, 2006), emphasizing the importance of system-level assessments of individual neuronal populations. Transcriptome analysis is now possible even at a single-cell level (Shapiro et al., 2013), and many genome-wide methods are being miniaturized (Buenrostro et al., 2013; Smallwood et al., 2014), making similar multidimensional approaches a possibility for even low abundance neuronal cells. We propose our integrated analysis of photoreceptor development (rods and S-cone-like) as a blueprint for cell-type-specific analysis of other neurons. Epigenomic landscape of adult rods and cones demonstrates unique profiles even in closely related neuronal subtypes (Mo et al., 2016). A comparison of rod and cone P28 transcriptome data (reported here) with that of Mo et al. (2016) revealed dissimilarities in photoreceptor gene expression profiles, which can be attributed to different methods of cell purification and genetic background of mouse lines. Further integration of transcriptome datasets with chromatin state of developing rods that we are generating and with the reported epigenomic data (Mo et al., 2016) would permit inference of functional interactions among the regulatory components in the omics landscape of photoreceptors and help in formulating testable hypotheses. Our studies also offer a framework for identifying convergent networks (Menche et al., 2015) for photoreceptor degeneration. The identification of a manageable number of common networks, upon which converge independent pathways affected by gene mutations, would facilitate the discovery of drug targets and therapies (Swaroop et al., 2010).

EXPERIMENTAL PROCEDURES

Isolation of Mouse Rod Photoreceptors

The Animal Care and Use Committee of the National Eye Institute (NEI) approved all procedures that involved mice. GFP-positive cells were purified from the retina of *Nrlp*-GFP and *Nrlp*-GFP;*Nrl*-KO mice (Akimoto et al., 2006). The retinas were dissected in Hank's Balanced Salt Solution (Life Technologies) at different postnatal stages of development, followed by dissociation in Accutase (Life Technologies) at 37°C for 10 min with constant agitation. After removing cell clumps by filtration, 4 mL PBS (Life Technologies) was added to dissociated cells. The cells were collected by centrifugation at 800 × *g* for 5 min and re-suspended in 1 mL PBS. GFP-positive cells were isolated by fluorescence-activated cell sorting (FACS) using FACS Aria II (Becton Dickinson) with a stringent precision setting to maximize the purity of the sorted cells. The purity of isolated GFP-positive cells was evaluated by re-sorting. After sorting, cells were processed with TRIzol LS (Invitrogen) following the manufacturer's instruction. For exon microarrays, *Nrlp*-GFP mouse retinas were treated in a similar manner, except that cells were dissociated by incubation in papain (Worthington Biochemical) supplemented with DNase I, superoxide dismutase, catalase, D- α -tocopherol acetate, and gentamycin at 28°C for 8 min. GFP-positive cells were collected into RNAProtect (QIAGEN).

Expression Data Generation and Analysis

Microarray data were generated using the WT-Ovation RNA Amplification System (NuGEN Technologies) and hybridized to GeneChip Mouse Exon 1.0 ST

Arrays (Affymetrix). Gene-level core RMA intensity values for each array were collected by Expression Console Software (Affymetrix) for mm8 annotation. Strand-specific RNA-seq data were generated using TruSeq RNA Sample Prep Kit-v2 (Illumina), as described previously (Brooks et al., 2012), and 76 base single-end reads were generated on Genome Analyzer IIx platform (Illumina). Transcript levels were quantified using Ensembl v73 transcriptome annotation, as described previously (Kaewkhaw et al., 2015).

Details of microarray hybridization, strand-specific RNA-seq, differential gene expression, RRBS DNA methylation, and bioinformatic analysis are provided in the [Supplemental Experimental Procedures](#).

In Situ Hybridization

C57BL/6 wild-type retina at P6 and P28 was fixed with 4% paraformaldehyde in PBS at 4°C overnight and transferred to 30% sucrose for overnight incubation at 4°C. The retina was then embedded in OCT mounting medium, cryosectioned at 12 μm thickness, and mounted on SuperFrost Plus glass slides. Retinal sections were then processed for fluorescent ISH using RNAscope technology (Advanced Cell Diagnostics), following the manufacturer's instructions with minor modifications. Probes were designed to recognize ~1,000 nt within each transcript (Table S3). Briefly, all probes were assayed using the RNAscope Multiplex Fluorescent kit. Slides were washed in PBS and incubated in Pretreat 2 solution at 100°C for 3 min, followed by incubation in Pretreat 3 solution at 40°C for 30 min. RNAscope probes were then hybridized at 40°C for 2 hr, followed by amplification steps according to the manufacturer's instructions. The signal was visualized and captured using a Zeiss 780 confocal microscope (Zeiss).

In Vivo Electroporation of shRNA

Custom small hairpin RNAs (shRNAs) were generated against the 3' UTR of the novel TCONS_00068375 gene according to the Addgene TRC cloning vector protocol (<https://www.addgene.org/tools/protocols/plko/#C>). Briefly, oligonucleotides containing the shRNA insert were annealed together, and the annealed oligonucleotides and the cloning vector (Addgene plasmid 10878) were digested with Agel-HF and EcoRI-HF. Digested oligonucleotides and gel-purified, digested plasmid were then ligated together. Constructed custom shRNA plasmids were transformed, amplified, and sequenced prior to in vivo electroporation. In vivo electroporation of the shRNA constructs were performed as described previously (Matsuda and Cepko, 2004). One eye of P0 CD1 mice was injected subretinally with 0.2 μL solution containing shRNA or control constructs and reporter plasmids, followed by five applications of 80-V pulses for 50 ms with a 950-ms interval to deliver DNA into the outer portion of the retina. To track total transfected cells as well as transfected rods, 500 ng/μL of each fluorescent reporter, CAGp-mCherry and *Nrlp*-GFP, respectively, was mixed with 2000 ng/μL of the control (Addgene plasmid 10879) or custom shRNA constructs. Two distinct shRNA constructs targeting the same gene and the control vector were injected in a single litter. Electroporated mice were sacrificed at P21, and the harvested retinas were fixed for 1 hr in 4% paraformaldehyde, cryopreserved in 30% sucrose, sectioned at 12 μm, and stained with DAPI. Fluorescence images were taken at 20× magnification on the Zeiss 700 confocal microscope.

Statistical Analysis

Statistical analysis of variances between two different experimental groups was conducted with Tukey's post hoc comparison test using SPSS. All experiments were repeated at least three times. The levels were considered significant at $p < 0.05$ (*), very significant at $p < 0.01$ (**), or not significant (n.s.).

ACCESSION NUMBERS

The accession numbers for the RNA-seq and microarray data reported in this paper are GEO: GSE74660 and GSE74657, respectively.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and three tables and can be found with this article online at <http://dx.doi.org/10.1016/j.celrep.2016.10.074>.

AUTHOR CONTRIBUTIONS

Overall Conceptualization, J.-W.K., H.-J.Y., N.G., and A.S.; Methodology and Investigation, J.-W.K., H.-J.Y., N.G., L.G., D.T.W., and R.V.; Transcriptome Data Analysis, M.J.B., V.C., J.J.B., A.W., A.B., and P.M.; Data Curation, M.J.B.; Network Analysis, V.C. and F.G.; De Novo Transcript Assembly and Alternate Splicing Analyses, G.K.; Database, K.D.K. and M.J.B.; Writing – Original Draft, H.-J.Y., J.-W.K., M.J.B., L.Z., A.S.; Writing – Review & Editing, J.-W.K., H.-J.Y., M.J.B., L.Z., G.K., V.C., T.C., and A.S.; Funding Acquisition, A.S.; Supervision and Project Administration, A.S.

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