Missense mutations in CRX homeodomain cause dominant retinopathies through two distinct mechanisms

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Yiqiao Zheng^{1,2}, Chi Sun², Xiaodong Zhang², Philip A. Ruzycki^{2,3} and Shiming Chen^{1,2,4}*

¹Molecular Genetic and Genomics Graduate program, Division of Biological and Biomedical Sciences, ²Department of Ophthalmology and Visual Sciences, ³Department of Genetics, ⁴Department of Developmental Biology, Washington University in St Louis, Saint Louis, Missouri, USA

*To whom the correspondence should be addressed. Mailing address: 660 South Euclid Avenue, MSC-8096-06-06, St. Louis, MO 63110, USA Tel. 314 747 4350 Fax. 314 747 4211 chenshiming@wustl.edu

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Summary

Homeodomain transcription factors (HD TFs) are instrumental to vertebrate development. Mutations in HD TFs have been linked to human diseases, but their pathogenic mechanisms remain elusive. Here we use *Cone-Rod Homeobox (CRX)* as a model to decipher the disease-causing mechanisms of two HD mutations, *p.E80A* and *p.K88N*, that produce severe dominant retinopathies. Through integrated analysis of molecular and functional evidence *in vitro* and in knock-in mouse models, we uncover two novel gain-of-function mechanisms: *p.E80A* increases CRX-mediated transactivation of canonical CRX target genes in developing photoreceptors; *p.K88N* alters CRX DNA-binding specificity resulting in binding at ectopic sites and severe perturbation of CRX target gene expression. Both mechanisms produce novel retinal morphological defects and hinder photoreceptor maturation distinct from loss-of-function models. This study reveals the distinct roles of E80 and K88 residues in CRX HD regulatory functions and emphasizes the importance of transcriptional precision in normal development.

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Introduction

Homeodomain transcription factors (HD TFs) play a fundamental role in vertebrate development. Members of the HD TF family are characterized by the presence of a highly conserved 60 amino acid helixturn-helix DNA-binding domain known as the homeodomain (HD). The HD is one of the most studied eukaryotic DNA-binding motifs since its discovery in *Drosophila* homeotic transformations^{1,2}. Hundreds of HD TFs have subsequently been documented in regulating gene expression programs important for body plan specification, pattern formation and cell fate determination^{1,2}. Mutations in HD TFs have been linked to many human diseases, including neuropsychiatric and neurodegenerative conditions^{3,4}. Although significant progress has been made in understanding HD-DNA interactions, uncovering the pathogenetic mechanisms of disease-causing missense mutations in HD have proven challenging.

The retina has long been used as a model system to study the role of HD TFs in normal central nervous system (CNS) development and in neurological diseases⁵. During retinogenesis, HD TFs play essential roles in the patterning of neuroepithelium, specification of retinal progenitors and differentiation of all retinal cell classes that derive from a common progenitor⁶. Importantly, many HD TFs are shared between the brain and the retina during development and mutations in these TFs can lead to disease manifestation in both tissues⁷⁻¹². The accessibility and wealth of available molecular tools makes the retina a valuable tool to decipher the pathogenic mechanisms of HD TF mutations associated with neurological diseases.

Here, we study CRX, a HD TF essential for photoreceptor cells in the retina, as a model to understand how single amino acid substitutions in the HD impact TF functions and cause blinding diseases. Photoreceptors are the most numerous neurons in the retina and are specialized to sense light and initiate vision through a process called phototransduction. Animal studies have demonstrated that *Crx* is first expressed in post-mitotic photoreceptor precursors and maintained throughout life^{13,14}. Loss of CRX results in impaired photoreceptor gene expression, failure of maturation and rapid degeneration of immature, non-functional photoreceptors¹⁵. Protein-coding sequence variants in human *CRX* have been associated with inherited retinal diseases (IRDs) that affect photoreceptors: Leber congenital amaurosis (LCA), cone-rod dystrophy (CoRD), and retinitis pigmentosa (RP) (OMIM:602225). However, the recessive phenotype observed in *Crx* knockout mouse models fails to recapitulate many dominant human *CRX* mutations that arise *de novo*¹⁵.

CRX contains two functional domains – the N-terminal HD and C-terminal activation domain (AD) (Figure 1A), both are required for proper activation of target genes and maintenance of normal *Crx* mRNA transcript abundance^{16,17}. To understand how CRX HD mutations cause diseases, we have previously reported a mutation knock-in mouse model carrying a hypomorphic mutation *p.R90W* (R90W) in CRX HD^{18,19}. We found that R90W mutation produces a recessive loss-of-function phenotype very similar to that of *Crx* knockout mice.

Intriguingly, several missense mutations within the same HD recognition helix as R90W, including p.E80A (E80A) and p.K88N (K88N), are linked to severe dominant IRDs²⁰⁻²² (Figures 1A and 1B). Here, we utilized a multi-omics approach to investigate the functional consequences of the E80A and K88N mutations on CRX regulatory activities and photoreceptor development (Figure S1). Comparison of the *in vitro* HD-DNA binding models of CRX and disease variants generated by Spec-seq revealed unique specificity changes of each mutant protein. Introduction of each mutation into the endogenous Crx locus generated knock-in mouse models Crx^{E80A} and Crx^{K88N} that reproduced dCoRD- and dLCA-like phenotypes. ChIP-seq analysis of CRX binding *in vivo* revealed mutation-specific changes in CRX targetome, consistent with mutation-specific DNA binding changes *in vitro*. Retinal RNA-seq analysis uncovered two distinct mechanisms by which the two HD missense mutation. Our results highlight the importance of residues E80 and K88 in CRX-mediated transcriptional regulation during photoreceptor development and the diverse mechanisms by which HD missense mutations can affect TF functions and lead to severe dominant neurological diseases.

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Results

K88N but not E80A mutation alters CRX HD DNA-binding specificity in vitro

CRX belongs to the *paired*-like HD TF family that recognize a 6-bp DNA motif in a stereotypic way²³⁻²⁹. Extensive studies of the HD have revealed a canonical HD-DNA recognition model where recognition of the 3' region (bases 4-6) of the HD DNA binding site is mediated by specificity determinants within the conserved HD recognition helix, corresponding to CRX residues 80-96^{23-26,29,30} (Figure 1B). In particular, HD residue 50, equivalent to CRX K88 residue (Figure 1A), is the specificity determinant in *paired*-like HD TF-DNA interactions^{24,25}. Since the three disease-associated HD missense mutations, E80A, K88N and R90W, are located within the CRX HD recognition helix, we wondered if these mutations change CRX HD DNA-binding specificity.

We adapted a high-throughput *in vitro* assay, Spec-seq, that determines protein-DNA-binding specificity by sequencing³¹⁻³³. Spec-seq was developed based on the traditional electrophoretic mobility shift assay (EMSA) to measure protein-DNA interactions. Spec-seq allows us to measure the relative binding affinities (i.e., specificity) for a library of HD binding motifs in parallel and generate quantitative binding models for different CRX HDs (Figures 1C). Based on the HD-DNA interaction model, we designed and tested a Spec-seq library containing all possible monomeric HD motifs (TAANNN) (Figure 1D).

We first obtained the wild-type (WT) CRX HD DNA binding model with Spec-seq using bacteriallyexpressed and affinity-purified HD peptides (Figures 1E-1G, STAR Methods). Relative binding energies of CRX WT HD from two experiments showed strong correlation (*r*: 0.984) and noise level (0.114 kT) within the expected range in typical Spec-seq data (Figure 1F). Binding energy model of WT HD was then generated by applying multiple linear regressions on the relative binding energies of all sequences within two base-pair mismatches to the WT CRX consensus (TAATCC)¹³ (Figure 1G, STAR Methods). A clear preference for CC bases at the 3' end of the motif is consistent with known CRX binding preference *in vitro* and *in vivo*^{34,35}.

We next sought to understand how disease mutations affect CRX DNA-binding specificity. We purified all mutant HD peptides following the same protocol as WT HD peptides and verified their DNA binding (Figures S2A-S2D)¹⁷. Comparison of the relative binding energies between each pair of mutant and WT HD revealed distinct effects (Figures 1H-1J). By definition, the consensus DNA binding motif of a testing peptide has a relative binding energy of 0kT. The relative binding energy difference between nucleotide variants and the consensus motif correlates with the DNA binding specificity of the testing peptide. We found that R90W HD and E80A HD both prefer the same consensus motif as WT (Figures 1K and 1L). R90W HD bound with slightly higher specificity than WT, as demonstrated by most data points falling above the diagonal line (Figures 1H and 1K), suggesting that R90W HD is more sensitive than WT to binding sequence variations. In contrast, when comparing E80A HD with WT HD, many data points fell below the diagonal line and the relative binding energies regressed towards 0 on the E80A axis (Figures 1I and 1L). This suggests that E80A HD bound with lower specificity than WT HD and thus was more tolerant to base variations in the HD DNA motif. Different from R90W and E80A, K88N mutation dramatically altered CRX HD DNA-binding specificity (r: 0.160) (Figures 1J and 1M). The K88N preferred binding sequence (TAAT/ATT/A) is referred to as N88 HD motif hereafter. K88N HD also had the largest degree of discrimination from its preferred to the weakest binding motif, suggesting that it is most sensitive to variants in the HD DNA motif. As a control, we tested a second library with the TAANNN sites on the reverse strand and obtained similar results (Figures S2E-S2L). Together, these results indicate that while E80A mutation does not affect CRX HD DNA binding specificity, the K88N mutation dramatically alters the specificity in vitro.

E80A protein binds to WT sites while K88N occupies novel genomic regions with N88 HD motifs in vivo

Next, we asked if changes in DNA-binding specificity affected mutant CRX chromatin binding in developing photoreceptors. We first created two human mutation knock-in mouse models, Crx^{E80A} and Crx^{K88N} , each carrying a single base substitution at the endogenous Crx locus (Figures S3A and S3B, STAR Methods). We confirmed that Crx mRNA was expressed at comparable levels in WT and mutant retinas (Figure S3C), and the full-length CRX proteins were readily detectable in the nuclear extracts from all samples (Figure S3D). We then obtained genome-wide binding profiles for each CRX variant by chromatin immunoprecipitation followed by sequencing (ChIP-seq) on mouse retinas at P14, a time when all retinal cell

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types are born, photoreceptor specification is completed in *WT* animals, and prior to any observed cell death in other CRX mutants previously characterized^{18,36}. To focus on changes specific to each mutant CRX protein, only homozygous animals were used for ChIP-seq profiling.

Unsupervised clustering of all CRX binding sites revealed two major clusters (Figure 2A, STAR Methods). Cluster 1 consisted of canonical WT CRX binding sites that are also occupied by CRX E80A protein (Figures 2A and 2B). Similar to WT CRX, CRX E80A binding *in vivo* was mostly enriched in intronic, followed by intergenic and TSS regions (Figure 3D). In contrast, CRX R90W, that also showed similar consensus preference to WT *in vitro*, failed to produce significant DNA binding *in vivo* (Figures 2A-2C, STAR Methods). This suggests that the retinopathy phenotype of $Crx^{R90W/W}$ is likely due to loss of binding at canonical WT CRX binding sites. Intriguingly, while CRX K88N showed loss of binding at canonical CRX binding sites, it gained a small set of binding sites (Cluster 2, Figures 2A-2C). *De novo* motif searching with DREME³⁷ under CRX peaks in each genotype revealed enrichment of monomeric HD motifs (Figure 2E) consistent with those found in Spec-seq (Figures 1G and 1K-1M), highlighted by a change in enriched HD motif from WT CRX HD type to N88 HD type in the $Crx^{K88N/N}$ retinas. Consistency with *in vitro* binding models suggests that *in vivo* changes in CRX chromatin binding were at least in part driven by the intrinsic changes in HD-DNA binding specificity by each individual mutation.

E80A and K88N mutations affected the expression of CRX-dependent activated genes in opposite directions in a critical time window of photoreceptor differentiation

To understand how different CRX mutations affected gene expression at canonical and *de novo* binding sites and how these changes impair photoreceptor differentiation, we turned to bulk RNA-seq analysis from the developing retinas at P10. At P10, photoreceptors have started to differentiate, and the expression of many photoreceptor genes undergo exponential increase^{38,39}. To focus on the most relevant expression changes, we first defined a set of genes that most likely depend on CRX activity nearby for expression (Figure S4A, STAR Methods). Briefly, we associated each CRX ChIP-seq peak to the nearest gene, filtered only genes with nearby CRX ChIP-seq peaks, and further narrowed the list of genes to those significantly down-regulated in the loss of function mutant $Crx^{R90W/W}$. Gene ontology (GO) analysis confirmed that this putative CRX-dependent gene set is associated with biological processes related to photoreceptor development and functions (Figure S4B). This set of putative CRX-dependent genes also showed developmental dependent gain in expression, consistent with CRX's primary function as a transcriptional activator (Figure S4C). As a control, CRX-independent genes were constitutively expressed and largely involved in general cellular processes (Figures S4D-S4F). Therefore, the CRX-dependent gene set comprises genes important for photoreceptor differentiation and functional maturation and are dependent on CRX for activation. We denote these genes as "CRX-dependent activated genes" (*CRX-DAGs*).

Next, we sought to understand how each mutation affected photoreceptor differentiation. One way of measuring the progression of photoreceptor differentiation is to determine the similarity in CRX-DAG expression in a given sample with that of known developmental ages in WT control animals. We thus performed sample-wise correlation of CRX-DAG expression obtained in our P10 samples with a previously published RNA-seq dataset of normal mouse retinal development (Figure 3A)³⁸. As expected, our P10 WT sample showed strong correlations with all developmental ages in the published WT control dataset. A stronger correlation with early ages (P3, P7, P10) and a weaker correlation with later ages (P14, P21) is also an indication of ongoing photoreceptor differentiation at P10. Unlike the WT sample, $Crx^{E80A/4}$ and $Crx^{E80A/4}$ samples both showed a stronger correlation with later developmental ages (P14, P21) but a weaker correlation with earlier postnatal ages (P3, P7). Since the CRX-DAGs are normally developmentally upregulated, this shift toward correlation with later developmental ages suggested that these genes were prematurely upregulated in the P10 Crx^{E80A} mutant retinas. In contrast, $Crx^{K88N/+}$ and $Crx^{K88N/+}$ samples both showed a weaker correlation with all developmental ages when compared with WT samples in our dataset. This suggests that early photoreceptor differentiation was compromised in both Crx^{K88N} mutants, consistent with their association with early-onset LCA²². Importantly, $Crx^{R90W/W}$, also associated with LCA-like phenotype^{18,21}, displayed strong correlation with earlier ages (P3, P7) similar to WT, but only showed moderate correlation with later ages (P14, P21). This suggests loss of CRX function at canonical binding sites does not affect the initiation of photoreceptor differentiation, but WT CRX activity at these sites is required to sustain differentiation. Since CRX-DAG expression was more severely affected in Crx^{K88N} mutants than in $Crx^{R90W/W}$, the photoreceptor differentiation

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deficits seen in the Crx^{K88N} mutants cannot be explained solely by the loss of regulatory activity at canonical CRX binding sites. Overall, our sample-wise correlation analysis with normal retinal development dataset suggests that E80A and K88N mutations affected the expression CRX-dependent activated genes in opposite directions, implicating novel and distinct pathogenic mechanisms from the loss-of-function R90W mutation.

Crx^{E80A} retinas show up-regulation of rod genes but down-regulation of cone genes, underlying CoRD-like phenotype

Upon closer examination, we noted that not all CRX-DAGs were up-regulated in Crx^{E80A} mutants (Figure 3B). Hierarchical clustering of all *CRX-DAGs* using expression changes revealed two major groups (STAR Methods). In aggregate, when compared to WT, Group 1 genes were up-regulated in Crx^{E80A} mutants while Group 2 genes were down-regulated. We noted genes indicative of the two photoreceptor subtypes, rods and cones, could partially define the two groups (Figures 3C and 3D). For example, $Esrb^{40}$ and Nrl^{41} in Group 1 are important regulators of rod differentiation. Other genes in Group 1 are components of the phototransduction cascade in rods, including Rcvrn⁴², Rho⁴³, Gnatl^{44,45}, Pde⁶g⁴⁶, Abca4^{47,48}, Gnb1⁴⁹, Rdh12⁵⁰, $Cngbl^{51}$, $Rpl^{52,53}$. Mis-regulations of many of these genes have been associated with diseases that affect rod development, function, and long-term survival. The increased activation of these genes likely underlies the stronger correlation with later developmental ages in $Crx^{E80A+/}$ and $Crx^{E80A//}$ retinas (Figure 3A). In contrast, Group 2 genes, many down-regulated in Crx^{E80A} mutants, were implicated in cone development and functions. For example, $Gnat2^{54,55}$, $Pde6c^{56}$, and $Pde6h^{57,58}$ all act in the cone phototransduction cascade. Mis-regulation of these genes has also been implicated in different retinal dystrophies that primarily affect cone photoreceptors. Comparison of ChIP-seq signal revealed that peaks associated with Group 2 genes showed lower occupancy compared to Group 1 genes in the $Crx^{E80A/A}$ retinas (Mann-Whitney U test *p-value*: 9.51e-07) while no difference was observed in WT retinas (Mann-Whitney U test p-value: 0.541), suggesting loss of CRX activity likely underlies the down-regulation of Group 2 genes in the Crx^{E80A} mutants (Figure S5A). Collectively, the selective down-regulation of cone genes in Group 2 may explain the CoRD-like phenotype in adult Crx^{E80A} mutant mice described later.

Additionally, we noticed that a subset of genes not affected in $Crx^{R90W/W}$ (CRX-independent genes) were also down-regulated in Crx^{E80A} mutants (Figures S5B-S5D). Among these genes were transcription regulators important for early photoreceptor development, such as $Ascl1^{59}$, Rax^{60} , $Sall3^{61}$, and $Pias3^{62}$. The down-regulation of these factors coincided with the up-regulation of mature rod genes in P10 Crx^{E80A} retinas, suggesting that the E80A mutation might hamper the proper timing of photoreceptor differentiation.

Crx^{K88N} retinas display greater reduction of rod and cone genes than the loss-of-function mutants

 Crx^{K88N} retinas had the most severe gene expression changes among all mutants with down-regulation of both Group 1 (rod) and Group 2 (cone) genes (Figures 3B- 3D). The heterozygous $Crx^{K88N/+}$ retina displayed a similar degree of expression reduction as homozygous $Crx^{R90W/W}$, consistent with its association with dLCA²². Given the normal phenotype of heterozygous loss-of-function mutants - $Crx^{+/-}$ and $Crx^{R90W/+18}$, these results suggested that mutant CRX K88N not only failed to activate WT target genes, but also functionally antagonized WT CRX regulatory activity in differentiating photoreceptors. This antagonism might be associated with ectopic CRX K88N activity when bound to regulatory regions with N88 HD DNA motifs (Figures 2A and 2E). In the absence of WT CRX, $Crx^{K88N/N}$ retina displayed a more severe expression reduction of CRX-DAG than $Crx^{R90W/W}$, raising the possibility that CRX K88N also antagonized the activity of other transcriptional regulators important for photoreceptor differentiation. Supporting this possibility, a set of CRX-independent genes were also misregulated in both heterozygous and homozygous Crx^{K88N} mutants, we noted that a number of these downregulated genes are also involved in photoreceptor functional development (Figures S5B, S5E, and S5F). Overall, CRX K88N is associated with greater gene expression changes than other CRX HD mutants, which is likely attributed to ectopic regulatory activity.

Both *E80A* and *K88N* mutants show compromised rod/cone terminal differentiation in young adults

Since $Crx^{+/-}$ and $Crx^{R90W/+}$ mutant mouse models showed a late-time recovery in photoreceptor gene expression and function¹⁹, we sought to determine the degree of photoreceptor differentiation in Crx^{E80A} and Crx^{K88N} mutants at P21. At this age, the normal retina has largely completed terminal differentiation with photoreceptor gene expression reaching a plateau⁶³. However, when P21 CRX HD mutant retinas were

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examined for the expression of genes under the GO term *detection of light stimulus* (GO:0009583), which comprises genes in both rod and cone phototransduction cascades, many genes failed to reach *WT* levels, despite variable degrees of impact across different HD mutants (Figure 4A). Crx^{E80A} mutants, in contrast to the increased rod gene expression at P10, displayed a deficiency in both cone and rod phototransduction genes at P21 (Figure 4B, Supplementary Table 6). This suggests that premature gene mutant CRX E80A transcriptional activity fails to sustain photoreceptor terminal differentiation and ultimately results in non-functional and severely affected photoreceptors. In comparison, Crx^{K88N} mutants showed severely reduced expression of rod/cone phototransduction genes at both P10 and P21 (Figures 3B-3D and 4B).

To assess retinal morphology and photoreceptor subtype-specific gene expression at the cellular level, we performed immunohistochemistry analysis on P21 retinal sections. In *WT* animals, a hallmark of photoreceptor maturation is the outgrowth of photoreceptor outer segments (OS) filled with proteins necessary for the phototransduction. We thus performed hematoxylin and eosin (H&E) staining on P21 sagittal retinal sections to visualize changes in retinal layer organization, focusing on photoreceptor layers – outer nuclear layer (ONL) and OS. Compared to the well-organized ONL in *WT* retinas (Figure 5A), all mutants showed variable degrees of ONL disorganization, forming waves, whorls, and rosettes (Figures 5B-5E). The ONL disorganization was more severe in homozygotes than in heterozygotes for both mutations and Crx^{K88N} mutants were more severely affected than Crx^{E80A} mutants. Photoreceptor OS layer was formed in the $Crx^{E80A/+}$ retinas, but absent in $Crx^{E80A/A}$, $Crx^{K88N/+}$, and $Crx^{K88N/N}$ mutant retinas. Inner retinal layers, including inner plexiform layer (IPL) and ganglion cell layer (GCL) were not as severely affected as the outer retinal layers, supporting a model that the mutant morphological abnormalities largely originated from the diseased photoreceptors. These morphological abnormalities were distinct from the degenerative phenotypes of other Crx mutant models reported previously^{18,64,65}.

Next, we sought to determine the expression of the rod-specific visual pigment rhodopsin (RHO) and cone arrestin (mCAR) in the P21 mouse retinas. In *WT* retinas, RHO is trafficked to the rod OS while mCAR is present in the cone OS and IS (inner segment), cell body, and synaptic terminals (Figure 5F). Unlike *WT* retina, all mutants lacked mCAR immunoreactivity (Figures 5G-5J), consistent with the loss of cone gene expression shown by RNAseq (Figure 4B). In $Crx^{E80A/+}$ retinas, RHO staining was localized to the OS layer; in $Crx^{E80A/A}$ and $Crx^{K88N/+}$ retinas, positive RHO staining was observed within the whorls and rosettes; in $Crx^{K88N/+}$ mutant retinas, RHO staining was completely absent. Importantly, we did not observe mis-localized RHO staining in the inner retinal layers suggesting that the developmental programs of other retinal cell types were not directly affected by E80A or K88N mutation. Overall, abnormalities in the cone/rod gene expression matched the corresponding human disease diagnosis^{20,22}, and the phenotypic severity correlated with the degree of mis-regulation of CRX target genes in the corresponding RNAseq dataset. Thus, these results support a model that CRX HD mutation-mediated mis-regulation of gene expression disrupts photoreceptor terminal differentiation and leads to defects in retinal layer organization and OS formation.

E80A and K88N mouse models show visual function deficits that recapitulate human diseases

To understand the consequences of disrupted photoreceptor differentiation on visual function, we measured electroretinogram (ERG) responses to light stimuli for *WT* and mutant mice at one month of age (Figures 5K-M). In response to incremental changes of light intensities, *WT* animals showed corresponding amplitude increases in dark adapted A-waves (rod signals) and B-waves (rod-evoked bipolar cell signals), as well as in light-adapted B-waves (cone-evoked bipolar cell signals). The three severe mutants, $Crx^{E80A/A}$, $Crx^{E80A/A}$, and $Crx^{E80A/A}$ had no detectable dark-adapted or light-adapted ERG responses, suggesting that these mice have no rod or cone function and are blind at young ages. The null ERG phenotype of the Crx^{K88N} animals is consistent with the clinical LCA phenotype in humans²². In contrast, $Crx^{E80A/+}$ animals retained partial rod ERG responses as indicated by the reduced A-wave and B-wave amplitudes (Figures 5K and 5L). Yet, $Crx^{E80A/+}$ animals had no detectable cone ERG responses, which is consistent with the CoRD clinical phenotype in humans²⁰ (Figure 5M). Taken together, the visual function impairment in each CRX HD mutant model, coincided with the morphological and molecular changes, suggesting that Crx^{E80A} and Crx^{K88N} mouse models recapitulate the corresponding human diseases.

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CRX E80A has increased transactivation activity and leads to precocious differentiation in *Crx*^{E80A} retinas

Lastly, we asked what might be the molecular mechanism that causes the mis-regulation of photoreceptor genes in the mutant retinas. Previous studies have established that reporter assays with the *rhodopsin* promoter in HEK293T cells can measure changes of CRX transactivation activity and inform the mechanisms by which photoreceptor genes are mis-regulated in CRX mutant retinas¹⁷. We thus tested the transactivation activity of the three CRX HD mutants on the *pRho-Luc* reporter in HEK293T cells (Figure 6A, STAR Methods). Consistent with published studies, R90W mutant had significantly reduced activity compared to WT CRX¹⁷. K88N mutant showed a similarly reduced activity as R90W, consistent with K88N's loss of binding at canonical CRX sites and failure to activate *CRX-DAGs in vivo* (Figures 2A, 2B, 3A-3C). In contrast, E80A mutant, which binds to canonical CRX sties, showed significantly increased transactivation activity on the *Rho* promoter. This hyperactivity of E80A protein at *Rho* promoter correlates with the upregulation of *CRX-DAGs* in the mutant retinas at P10 (Figures 3A-3C).

To determine the consequences of E80A hyperactivity on photoreceptor differentiation timing, we compared RHO protein expression during early postnatal retinal development using retinal section immunostaining (Figures 6B-6D). In *WT* retinas, most rods were born by P3 but had not differentiated (Figure 6B). Previously studies showed that RHO proteins were detected by IHC starting around P7 in *WT* retinas⁶³. In comparison, both $Crx^{E80A/+}$ and $Crx^{E80A/A}$ retinas showed positive RHO staining at P3 (Figures 6C and 6D). RHO⁺ cells were largely seen in the outer portion of the ONBL layer in $Crx^{E80A/+}$ retinas, and strikingly spread throughout the large presumptive ONL layer in $Crx^{E80A/A}$ retinas. The detection of RHO protein in P3 Crx^{E80A} mutant retinas indicates that photoreceptor differentiation program was precociously activated. Taken together, our results support a model that *E80A* and *K88N* mutations each perturbs CRX regulatory activity in a unique way, causes photoreceptor differentiation defects, and ultimately leads to distinct dominant disease phenotypes that recapitulate human diseases.

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Discussion

Through molecular characterization of mutant proteins, transcriptome and cellular profiling of developing mutant mouse retinas and ERG testing of adult retinas, we have identified two novel pathogenic mechanisms of CRX HD mutations, E80A and K88N, that are associated with dominant CoRD and dominant LCA in human^{20,21}. Distinct from the previously characterized loss-of-function R90W mutation^{18,19}, E80A and K88N mutations produce altered CRX proteins with gain of regulatory functions - CRX E80A is associated with increased transcriptional activity and CRX K88N has altered DNA binding specificity. Both CRX E80A and CRX K88N proteins impair photoreceptor gene expression, development and produce structural and functional deficits in knock-in mouse models, recapitulating human diseases⁶⁴ (Figure 7). Thus, precise regulatory activity and target specificity of CRX are both essential for proper photoreceptor development and functional maturation.

The Crx^{E80A} and Crx^{K88N} knock-in mouse models recapitulate dominant human CRX-linked retinopathies and present novel morphological deficits. Published Crx mutant mouse models have established that decreased expression of CRX or loss of CRX transcriptional activity results in incomplete photoreceptor differentiation and ultimately rapid degeneration of immature photoreceptors^{15,18}. Despite the difference in molecular mechanisms, both Crx^{E80A} and Crx^{K88N} mouse models show defective photoreceptor terminal differentiation and develop whorls and rosettes in the ONL layers by P21, which are not observed in degenerative CRX mouse models¹⁸. No significant increase in cell death is observed in Crx^{E80A} and Crx^{K88N} mutant mouse retinas at P21, suggesting distinct pathogenic mechanisms than the photoreceptor degeneration previously reported in other CRX mouse models. P10 RNAseq analysis on cell markers of other neuronal cell types and Muller glia cells did not detect significant changes, suggesting that the developmental programs for these cell types were unaffected. Further studies are needed to systematically characterize changes in photoreceptor differentiation and retinal morphology during early development, and interactions between diseased photoreceptors and other major cell types in Crx^{E80A} and Crx^{K88N} mutant mouse retinas.

Precisely tuned CRX-mediated transcriptional activity is important to ensure proper photoreceptor development. Previous studies have demonstrated that CRX is activated shortly after cell cycle exit in retinal progenitor cells fated to be photoreceptors^{66,67}. CRX then works with other cell type specific TFs, such as NRL and NR2E3 (rods) or RORB and THRB (cones), to regulate gene expression important for cell-type specific structures and functions and drive the differentiation of photoreceptor precursors^{68,69}. Loss-offunction animal studies have established that CRX is primarily a transcriptional activator since loss of CRX protein or regulatory function is associated with decreased expression of photoreceptor-specific genes regulated by CRX¹⁹. Here, we find that CRX E80A mutant, that displays increased CRX transcriptional activator activity, is also deleterious to photoreceptor development. CRX E80A maintains WT DNA-binding specificity in vitro and binds to canonical CRX sites in mutation knock-in mouse retinas. Consistent with CRX E80A's hyper-activity at canonical CRX target *pRho*, in differentiating $Crx^{E80A/+}$ and $Crx^{E80A/A}$ retinas, many of the CRX-dependent activated genes implicated in rod functions are prematurely up-regulated at early postnatal ages. The RHO protein is also detected in developing rod photoreceptor in Crx^{E80A/+} and Crx^{E80A/A} retinas as early as P3. However, at later ages, rod photoreceptor terminal differentiation is impaired, resulting in disrupted retinal morphology and defective visual functions. Photoreceptor differentiation is precisely programmed via sequential and concerted gene expression programs within a defined time window^{66,67}. One explanation for adult Crx^{E80A} phenotypes is that CRX E80A associated hyper-activity precociously activates later stage genes in the absence of proper nuclear context and/or subcellular structures, which in turn negatively impacts early events in photoreceptor differentiation. Multiple possibilities could account for CRX E80A associated hyper-activity: (1) E80A mutation increases CRX interactions with its binding sites; (2) E80A mutation increases CRX interactions with the transcription complex or other TFs; (3) certain protein factors that could promote photoreceptor gene expression are up-regulated in CRX E80A expressing cells. While these possibilities are not mutually exclusive, evidence from the current study favors the first possibility since E80A HD is less discriminative of HD motif variants than WT HD as shown by Spec-seq (Figure 1I). Regulatory sequences of many photoreceptor genes contain both consensus and non-consensus CRX motifs^{13,14,17,34}. CRX E80A might interact better with non-consensus CRX motifs within binding sites, resulting in increased transcriptional output of target genes. Large scale reporter assays (e.g. Massively Parallel Reporter Assays) in the appropriate cellular context - neonatal mouse retinas - would be informative to understanding CRX E80A hyper-activity mechanism and the direct impacts of its hyper-activity.

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Although a global increase in expression is expected in Crx^{E80A} models, a subset of CRX dependent activated genes implicated in cone photoreceptor development and functions is down-regulated in both differentiating and mature mutant retinas. Cones are born prenatally in mice within an earlier time window than rods. Yet, cones undergo terminal differentiation in a similar postnatal window as rods to develop into subtypes of cones – M-cones v.s. S-cones. One model is that CRX E80A improperly activates later stage cone genes at a much earlier time window, disrupting cone terminal differentiation. Supporting this model, similar to the presented pRho data, CRX E80A also hyper-activates the S-cone opsin promoter (data not shown). Detecting premature activation of cone genes in Crx^{E80A} retinas will require analysis at early developmental ages. An alternative model for lack of cone markers in Crx^{E80A} retinas is that cones might be more sensitive to perturbations in CRX transcriptional activity, which impacts cone survival. It is known that cones depend on a different repertoire of TFs than rods for subtype terminal differentiation⁵⁹. It is possible that cone TFs respond differently to mutant CRX E80A hyper-activity, leading to the distinct expression changes in Crx^{E80A} mutant retains. Since cones only make up a very small portion of photoreceptors in mouse retinas⁷⁰, characterization of CRX E80A activity in purified cone population at earlier ages will be necessary to understand its selective effect on the cone differentiation program and help to elucidate WT CRX regulatory principles in early photoreceptor development. Collectively, CRX transcriptional activity at its canonical genomic binding sites need to be maintained within a quantitatively defined range such that CRX dependent target genes are activated in a precise temporal pattern to ensure proper photoreceptor differentiation.

CRX DNA binding specificity constitutes an additional regulatory program required for the fidelity of photoreceptor gene expression during development. Previous work has shown that subtle differences in HD motif content could have distinct transcriptional and functional outcomes. For instance, studies in D. melanogaster demonstrate that the sequential use of HD DNA motif variants defines different classes of enhancers and coordinates the precise timing of gene expression patterns during embryonic development⁷¹. In our current study, the K88N mutation alters CRX DNA-binding specificity that redirects CRX binding to binding sites with N88 HD motifs in the mutant mouse retinas. Crx^{K88N} retinas show more severe perturbations in photoreceptor gene expression and development than in loss-of-function mutant $Crx^{R90W/W}$. Concomitantly, $Crx^{K88N/+}$ and $Crx^{K88N/N}$ mice display the most severe photoreceptor morphological deficits observed in any Crx mouse models and show absence of visual functions in young adults. In developing WT mouse retinas, the HD motif preferred by HD TFs with a glutamine (Q) at HD50 position encodes quantitatively different activity than the CRX consensus motif suggesting functional difference between HD binding site variants^{60,72}. It is likely that the severe Crx^{K88N} phenotypes are attributed to both loss of activity at canonical CRX motifs and ectopic binding and transcriptional activity of CRX K88N proteins at N88 HD motifs. Additionally, since a functional WT copy of CRX is retained in Crx^{K88N/+} retinas and CRX K88N prefers different motifs than WT CRX, the lack of WT activity alone cannot explain the severe expression and developmental deficits. Alternatively, these results suggest involvement of additional regulatory mechanisms: CRX K88N activity at N88 HD motifs might (1) ectopically activate genes whose expression prevents the progression of development or inactivate genes required for development; (2) interfere with other HD TFs that also recognize N88 HD motifs; (3) lead to epigenetic alterations that antagonize normal CRX functions. Many other HD containing TFs are expressed in developing mouse retina, including OTX2, RAX, VSX2, PAX6, SIX3/6, and LHX family⁶. Different from CRX, these HD TFs are essential for gene regulation in retinal progenitor cells and/or in other retinal cell lineages. It is very likely that CRX correctly targets photoreceptor genes at least through its DNA binding specificity and alteration of CRX DNA binding specificity would mis-regulate genes originally targeted by other HD TFs and lead to perturbations in GRNs and development. To date, most studies have focused on CRX activity at cis-regulatory sequences enriched for the WT CRX consensus motifs^{73,74}. Systematic comparison of transcriptional activity at N88 HD motifs and WT consensus motifs in the context of photoreceptor development in both WT and mutant retinas would be needed to substantiate the impact of mutant CRX K88N activity at different HD motifs. These experiments will also help clarify the pathogenic mechanisms in the Crx^{K88N} models and extend our knowledge of CRX regulatory grammar and activation of photoreceptor genes during development.

One limitation of this work is that effects of E80A and K88N mutations on CRX HD-DNA interactions have been evaluated at monomeric HD motifs and with homogenous protein species both *in vitro* and *in vivo*. Further evaluation of WT and mutant CRX binding at dimeric motifs will be desirable, since selected dimeric HD motifs are known to mediate HD TF cooperativity to ensure gene expression fidelity

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during development^{75,76}. Relatedly, we also need to address how CRX WT and mutant E80A or K88N proteins interact at HD binding motifs – whether they cooperate or compete with each other, whether these interactions are HD motif sequence-dependent, and how gene expression is impacted by CRX cooperativity or competition. While CRX HD mediates both TF-DNA interactions and protein-protein interactions, evaluation of how E80A and K88N mutations impact CRX interaction with other important photoreceptor TFs and how perturbations in these interactions lead to disease phenotypes warrant further study.

Since the spatial structures and HD-DNA contact models of HD proteins are evolutionarily conserved, our study of CRX provides valuable molecular insights for HD mutations implicated in other diseases. For example, *p.E79K* substitution (corresponds to CRX E80) in OTX2 HD is associated with dominant early-onset retinal dystrophy in two Caucasian Canadian families⁷⁷, heterozygous *p.R89G* (corresponds to CRX R90) mutation in OTX2 HD causes severe ocular malformations⁷⁸, and missense mutations of the CRX K88 and R90 homologous residues in PITX2 HD are associated with Rieger syndrome⁷⁹. It is likely that these mutations affect HD activity in similar ways as observed in CRX, and the exact disease manifestation is determined by cell-type specific mechanisms. The retina is readily accessible and a broad range of molecular tools are available for *ex vivo* and *in vivo* manipulations. We believe that CRX is an ideal model to understand the pathogenic mechanisms of HD mutations and to test therapeutic regimens, which would ultimately benefit the study of HD TFs and their associated diseases in other tissues and organs.

Collectively, our findings support a unifying model in which precise CRX interaction with cisregulatory sequences is essential for gene expression and functional maturation during photoreceptor development. Mutations that perturb CRX target specificity or transactivation activity impair the integrity of CRX-mediated transcriptional regulation and result in distinct morphological and functional deficits (Figures 7A-7D). Regulatory activity at ectopic N88 HD motifs antagonizes normal CRX functions and leads to aberrant photoreceptor development and early vision loss. This model also emphasizes the importance of tailoring gene therapy regimens to tackle individual pathogenic mechanisms. For instance, while supplementing WT CRX might be sufficient to rescue a hypomorphic/loss-of-function mutant, simultaneous elimination of a gain-of-function *CRX* product would be necessary to rescue dominant mutants, as exemplified in a recent report of allele-specific gene editing to rescue dominant CRX-associated LCA7 phenotypes in a retinal organoid model⁸⁰. We believe that this principle also applies to other dominant neurological diseases. Additionally, with the refinement of the CRX mechanistic model, when new disease mutations are identified, genetic counsellors can now provide more informed predictions of disease progression and future visual deficits. This information is important for individuals to be psychologically prepared and seek necessary assistance to improve their quality of life.

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Author Contributions

S. Chen conceived and supervised the study. S. Chen, P. Ruzycki, C. Sun and Y. Zheng designed the experiments. Y. Zheng performed Spec-seq and luciferase experiments; X. Zhang performed CRX ChIP-seq experiments; C. Sun performed immunochemistry and ERG experiments. Y. Zheng analyzed Spec-seq, CRX ChIP-seq, RNA-seq, and luciferase data; P. Ruzycki assisted in CRX ChIP-seq and RNA-seq data analysis; C. Sun analyzed immunochemistry and ERG data. Y. Zheng wrote the original draft; S. Chen, P. Ruzycki, C. Sun, and Y. Zheng revised the manuscript. All authors read and approved the final manuscript.

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Declaration of Interests

The authors declare no competing interests.

Inclusion and diversity

We support inclusive, diverse, and equitable conduct of research.

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Main figure titles and legends

Figure 1. Disease associated missense mutations altered CRX HD DNA binding specificity. (A) Diagram of CRX functional domains: Homeodomain (HD) for DNA-binding and Activation Domain (AD) for target gene transactivation. The three missense mutations in this study are located at the C-terminus of CRX HD and associated with different retinal diseases in human. Number in the parenthesis denotes the CRX HD position of the corresponding mutated residue. (B) Alignments of HD recognition helix sequences for the indicated HD proteins for which HD missense mutations have been associated with inherited diseases. Accession numbers can be found in Supplementary Table S1. Missense variants in this study (highlighted) are located at highly conserved residues across species and different HD TFs. (C) Spec-seq experimental workflow (STAR Methods). (D) Spec-seq library design of monomeric HD binding sites. (E) EMSA gel images of Spec-seq experiments with different CRX HD species. B_x: Bound. B.: Unbound. (F) Relative binding energy comparison from two different experiments with WT HD. (G) Binding energy model for WT CRX HD. (H-J) Relative binding energy comparison between WT HD and R90W HD (H), E80A HD (I), or K88N HD (J). Consensus sequence is defined to have relative binding energy of 0kT (TAATCC for WT, R90W and E80A, TAATTA for K88N). The diagonal line of identity is represented in grey dash. The orange dashed line shows the best linear fit to the data. (K-M) Binding energy models for R90W HD (K), E80A HD (L), and K88N HD (M). Only sequence variants within two mismatches to the corresponding consensus sequences were used to generate binding models. Negative binding energy is plotted such that bases above the x-axis are preferred bases and bases below the x-axis are unfavorable bases. Constant bases (TAA) carried no information are drawn at arbitrary height in grey.

Figure 2. CRX E80A binds to WT sites while CRX K88N occupies novel genomic regions enriched for N88 HD motif *in vivo*. (A) Enrichment heat map depicting CRX ChIP-seq normalized reads centered at all possible CRX peaks ± 2kb, sorted by binding intensity in *WT* samples. Clusters were defined by hierarchical clustering of CRX binding intensity matrix from all genotypes (STAR Methods). (B-C) Genome browser representations of ChIP-seq normalized reads for different CRX species in P14 *WT* and mutant mouse retinas at *Rho* and *Atf2*. (D) Enrichment heatmap showing fraction of CRX ChIP-seq peaks fall in different genomic environments. (E) Logo representations of *de novo* found short HD motifs under CRX ChIP-seq peaks in *WT* and mutant mouse retinas with DREME *E-value* on the right.

Figure 3. CRX-dependent activated genes affected in opposite directions in developing *Crx*^{E80A} **and** *Crx*^{K88N} **mutant retinas** (A) Heat map showing sample-wise Pearson correlations of the expression of all CRX-dependent activated genes between P10 *WT* and HD mutant mouse retinas in this study (rows) with post-natal *WT* retinas from age P3 to P21 (columns, data from GSE87064). (B) Heat map showing the expression changes of DEGs in CRX-dependent activated gene set in HD mutant mouse retinas at P10. (C-D) Heat maps showing expression changes of selected photoreceptor genes from Group 1 and Group 2. Color scale identical to (B).

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Figure 4. Photoreceptor genes important for phototransduction are down-regulated in all HD mutants

(A) Box plot showing that genes in the detection of light stimulus GO term were down-regulated and affected to various degrees in different adult (P21) HD mutant mouse retinas. (B) Heat map showing that expression of both cone and rod phototransduction genes were down-regulated in adult (P21) HD mutant mouse retinas. Annotation of rod and cone enrichment of each gene is in Supplementary Table 6. See Supplementary Figure S6 for the developmental expression dynamics of these genes.

Figure 5. Only $Crx^{E80A/+}$ retinas maintain photoreceptor OS and residual rod ERG response (A-E) Hematoxylin-eosin (H&E) staining of P21 retina sections show that photoreceptor OS layer is absent in all mutant retinas except $Crx^{E80A/+}$. OS: outer segment; ONL: outer nuclear layer; INL: inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell layer. Scale bar, 100µm. (F-J) Rhodopsin (RHO, red) immunostaining is present in $Crx^{E80A/+}$, $Crx^{E80A/A}$, and $Crx^{K88N/+}$ retinas and absent in $Crx^{K88N/N}$ retina. Cone arrestin (mCAR, green) immunostaining is absent in all mutant retinas. Nuclei were visualized by DAPI staining (Blue). Scale bar, 100µm. (K-M) The electroretinogram responses (ERG) recorded from 1-month mice. Error bars represent the standard error of the mean (SEM, $n \ge 4$). *p-value*: Two-way ANOVA and Tukey's multiple comparisons. ****: $p \le 0.0001$. ns: >0.05.

Figure 6. CRX E80A hyperactivity underlies precocious photoreceptor differentiation in Crx^{E80A} retinas. (A) Boxplot showing luciferase reporter activities of different CRX variants. *P-values* for one-way ANOVA with Turkey honestly significant difference (HSD) test are indicated. *p-value*: ****: ≤ 0.0001 , ***: ≤ 0.001 , ns: >0.05. (B-D) Rhodopsin (RHO, green) immunostaining is absent in P3 *WT* retina but detected in $Crx^{E80A/+}$ and $Crx^{E80A/+}$ retinas. Nuclei are visualized by DAPI staining (Blue). Arrow indicates the sporadic RHO staining in $Crx^{E80A/+}$ sample. ONBL: outer neuroblast layer; GCL: ganglion cell layer. Scale bar, 100µm.

Figure 7. Missense mutations in CRX HD affect photoreceptor gene expression and leads to distinct retinal disease phenotypes through gain- and loss-of-function mechanisms.

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STAR Methods

Resource Availability

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact Shiming Chen (chenshiming@wustl.edu).

Materials Availability

• All unique/stable reagents generated in this study are available from the lead contact with a completed materials transfer agreement.

Data and Code Availability

- The raw sequencing data and processed data generated in this study have been deposited at NCBI.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

Animal study and sample collection

Ethics statement

All procedures involving mice were approved by the Animal Studies Committee of Washington University in St. Louis and performed under Protocol 21-0414 (to SC). Experiments were carried out in strict accordance with recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health (Bethesda, MD), the Washington University Policy on the Use of Animals in Research; and the Guidelines for the Use of Animals in Visual Research of the Association for Research in Ophthalmology and Visual Sciences. Every effort was made to minimize the animals' suffering, anxiety and discomfort.

Mutation knock-in mouse model generation

CRISPR/Cas9 based genome editing was performed to generate the Crx^{E80A} and Crx^{K88N} mice as previously described⁸¹. The Cas9 guide RNAs (gRNA) were designed based on proximity to the target amino acid and was synthesized using the MEGAshortscriptTM T7 Transcription Kit (Thermo Fisher Scientific, Waltham, MA). The gRNAs were subsequently tested for cutting efficiency in cell culture by the Washington University Genome Engineering and iPSC Center. The validated gRNA and Cas9 protein were then microinjected into the pronuclei of C57BI/6J- 0.5-dpc (days post coitum) zygotes along with the donor DNA, a 190-bp single stranded oligodeoxynucleotide (ssODN) carrying either the *c.239A*>*G* substitution for *p.E80A* mutation or the *c.264G*>*T* substitution for *p.K88N* mutation⁸². Embryos were then transferred into the oviduct of pseudo-pregnant female. Pups were generally delivered ~20 days after microinjection. Tissues from 10-day postnatal (P10) pups were collected by toe biopsy/tail for identification of the targeted allele by restriction digest (HinfI) of PCR amplified DNA first and then confirmed by Sanger sequencing (Genewiz).

Founders carrying the correct alleles were then bred with wild-type C57BL/6J mice (Jackson Laboratories, Bar Harbor, ME, Strain #000664) to confirm transmission. All experimental animals used were backcrossed at least 10 generations. Genotyping of mutation knock-in mice follows cycling conditions: 95°C for 2 min, 94°C for 30 sec, 60°C for 30 sec, 68°C for 60 sec, repeat steps 2-4 for 34 cycles, 68°C for 7 min, and hold at 4°C. After PCR reaction, the amplified DNA fragments wee digested with HinfI. Sequences of gRNAs, ssODNs and genotyping primers can be found in Supplementary Table S3. A representative DNA gel of HinfI digested genotyping DNA fragments can be found at Figure S3B.

RNA-seq sample collection and library preparation

For each genotype, three biological replicates, two retinas per replicate from one male and one female mouse were analyzed. All retinas were processed for RNA simultaneously using TRIzolTM Reagent (Invitrogen, Waltham, MA) following the manufacturer's protocol. The quantity and quality of the RNA was assayed

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using Bioanalyzer (Agilent, Santa Clara, CA). Samples with a minimum RNA integrity number (RIN) score of 8.0 were then selected for library construction as previously described¹⁹.

Chromatin immunoprecipitation and library preparation

CRX chromatin immunoprecipitation was performed as previously published⁸³. Briefly, pooled nuclear extracts from 6 retinae were cross-linked with formaldehyde prior to immunoprecipitation with anti-CRX antibody A-9 (#sc-377138, Santa Cruz Biotechnology, Dallas, TX). Input controls were included as background. The libraries were prepared following the standard ChIP-seq protocol⁸⁴. The quantity and quality of the ChIP-seq libraries was assayed using Bioanalyzer (Agilent, Santa Clara, CA) prior to sequencing.

ERG and Statistical Analyses

ERGs were performed on 1-month-old mice using UTAS-E3000 Visual Electrodiagnostic System (LKC Technologies Inc., MD). Mice were dark-adapted overnight prior to the tests. Mouse body temperature was kept at 37 ± 0.5 °C during the tests. Pupils were dilated with 1% atropine sulfate solution (Bausch and Lomb). Platinum 2.0 mm loop electrodes were placed on the cornea of each eye. A reference electrode was inserted under the skin of the mouse's head and a ground electrode was placed under the skin near mouse's tail. Retinal response to full-field light flashes (10 µs) of increasing intensity were recorded; maximum flash intensity for dark-adapted testing was 0.895 cd*s/m2. Following dark adapted tests, mice were light-adapted under light condition (about 29.2 cd/mm) for 10 minutes and exposed to 10 µs light flashes of increasing intensity; maximum flash intensity for light-adapted testing was 2.672 cd*s/m2. ERG responses of biological replicates were recorded, averaged and analyzed using Graphpad Prism 8 (GraphPad Software, CA). The mean peak amplitudes of dark-adapted A and B waves and light-adapted B waves were plotted against log values of light intensities (cd*s/m2). The statistics were obtained by two-way ANOVA with multiple pairwise comparisons (Tukey's).

Histology and Immunohistology chemistry

Enucleated eyes were fixed at 4°C overnight for paraffin embedded sections. Each retinal cross-section was cut 5 microns thick on a microtome. Hematoxylin and Eosin (H&E) staining was performed to examine retinal morphology. For IHC staining, sections firstly went through antigen retrieval with citrate buffer, and blocked with a blocking buffer of 5% donkey serum, 1% BSA, 0.1% Triton X-100 in 1x PBS (pH-7.4) for 1 h. Sections were then incubated with primary antibodies at 4°C overnight. Sections were washed with 1x PBS containing 0.01% Triton X-100 (PBST) for 30 min, and then incubated with specific secondary antibodies for 1 h. Primary and secondary antibodies were applied with optimal dilution ratios. All slides were mounted with VECTASHIELD® HardSetTM Antifade Mounting Medium with DAPI (Vector Laboratories, Inc., CA). Anti-Rho antibody was obtained from MilliporeSigma, O4886. Anti-mCAR antibody was obtained from MilliporeSigma, AB15282. All images were taken on a Leica DB5500 microscope. All images were acquired at 1000 μ m from ONH for \geq P21 samples and at 500 μ m from ONH for P3, P10 samples.

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Biochemistry

Protein expression

Expression plasmids for GST-WT, E80A, and R90W HDs were published previously¹⁷. Plasmid for GST-K88N HD was generated by site-directed mutagenesis from the pGEX4T2-CRX WT HD backbone. In vivo protein expression and purification was done as previously described¹⁷. Briefly, 0.05mM IPTG was added to *E. coli* BL-21 (DE3) cell cultures containing different CRX HD constructs at $OD_{600} = 0.6$. The cultures were incubated for 2 hours or until $OD_{600} = 2.0$ at 34°C and the cells were collected by centrifugation at 6000 rpm and 4°C for 15 minutes. Cell pellets were resuspended in 1x PBS (Corning, Corning, NY) and then lysed by sonication. 5mM DTT (Bio-Rad Laboratories, Inc., Hercules, CA) and 1% Triton X-100 (MilliporeSigma, Burlington, MA) was then added, and the mixtures were incubated with gentle shaking at 4°C for 30 minutes to maximize protein extraction. The separation of proteins from the cellular debris were then performed by centrifugation at 15 000 rpm for 10 minutes and filtered through a 0.45µm membrane. Glutathione Sepharose 4B resin (Cytiva, Marlborough, MA) was first equilibrated with PBS before adding to the supernatant. 5x HaltTM Protease Inhibitor Cocktail and PMSF was added to minimize degradation. The mixtures were incubated with gentle shaking at 4°C overnight before loading on GST SpintrapTM columns (Cytiva, Marlborough, MA). The protein was eluted following the manufacturer's protocol and buffer exchanged into CRX binding buffer⁸⁵ using Amicon centrifugal filters (MilliporeSigma, Burlington, MA). The protein stock was supplemented with 10% glycerol before aliquoted and stored at -80°C.

Protein quantification and visualization

The size and integrity of purified GST-CRX HDs were visualized with a native 12% Tris-Glycine SDS-PAGE gel in the absence any reducing agent. Protein concentration was measured by NanoDrop One^o Microvolume UV-Vis Spectrophotometers (ThermoFisher Scientific, Waltham, MA) and calculated using the equation: $C = (1.55 * A_{280}) - (0.76 * A_{260})$, where C is the concentration of the protein in mg/ml, A_{280} and A_{260} is the absorbance of protein samples at 280nm and 260nm respectively⁸⁶. The protein concentration obtained with this method was comparable with BCA protein quantification assays.

Spec-seq library synthesis and purification

Single-stranded Spec-seq library templates and IRDye 700-labeled reverse complement primers (Supplementary Table S1) were ordered directly from Integrated DNA Technologies (IDT, Coralville, Iowa). The synthesis and purification of the double-stranded libraries followed previously published protocols^{33,86}. Briefly, 100 pmol of template oligos and 125 pmol IRDye 700-labeled reverse complement primer F1 were mixed in Phusion® High-Fidelity PCR Master Mix (NEB, Ipswich, MA). A 15s denaturing at 95°C following a 10-minute extension at 52°C afforded duplex DNAs. Subsequently, the mixture was treated with Exonuclease I (NEB, Ipswich, MA) to remove excess ssDNA. The libraries were purified by MinElute PCR Purification Kit (QIAGEN, Hilden, Germany) and eluted in molecular biology graded water (Corning, Corning, NY).

EMSA and sample preparation for sequencing

The protein-DNA binding reactions was done in 1x CRX binding buffer (60mM KCl, 25mM HEPES, 5% glycerol, 1mM DTT)⁸⁵. A fixed amount (Supplementary Table S1) of IRDye-labelled DNA libraries were incubated on ice for 30 minutes with varying concentrations of wild type or mutant peptides in 20µl reaction volume. The reaction mixtures were run at 4°C in native 12% Tris-Glycine PAGE gel at 160V for 40min. The IRDye-labeled DNA fragments in the bound and unbound fractions were visualized by Odyssey® CLx and Fc Imaging Systems (LI-COR, Inc., Lincoln, NE). The visible bands were excised from the gels and DNAs were extracted with acrylamide extraction buffer (100mM NH₄OAc, 10mM Mg(OAc)₂, 0.1% SDS) then purified with MinElute PCR Purification Kit (QIAGEN, Hilden, Germany). The DNAs were amplified, barcoded by indexed Illumina primers. All indexed libraries were then pooled and sequenced on a single 1x50bp Miseq run

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at DNA Sequencing Innovation Lab at the Center for Genome Sciences & Systems Biology (CGS&SB, WashU).

<u>qRT-PCR</u>

For each replicate, RNA from 2 retinae of a mouse was extracted using the NucleoSpin® RNA kits (Takara Bio USA, Inc., San Jose, CA). RNA sample concentration and quality was determined with NanoDrop One^c Microvolume UV-Vis Spectrophotometers (ThermoFisher Scientific, Waltham, MA). 1 μg of RNA was used to synthesize cDNA with iScript cDNA Synthesis Kits (BioRad, Hercules, CA). Primers used in this study were listed in Supplementary Table S2. qRT-PCR reactions were assembled using SsoFastTM EvaGreen® Supermix with Low ROX following manufacturer's protocol (Bio-Rad Laboratories, Inc., Hercules, CA). Data was obtained from Bio-Rad CFX96 Thermal Cycler following a three-step protocol: 1 cycle of 95°C 3 min, 40 cycles of 95°C 10 sec and 60°C 30 sec. Data was exported and further processed with customized python script.

Western blot

Experiments were performed using two biological replicates with two retinas for each replicate. Nuclear extracts were prepared using the NE-PER Nuclear and Cytoplasmic Extraction Reagents (Thermo Fisher Scientific, Waltham, MA) following manufacturer's instructions. 1x Roche cOmplete[™], Mini Protease Inhibitor Cocktail (MilliporeSigma, Burlington, MA) was supplemented in all extraction reagents. 5mM of DTT was added immediately before sample denaturing and protein was separated by running on Invitrogen NuPAGE[™] Novex 4-12% Bis-Tris MiniGels (Invitrogen, Waltham, MA). Membrane transfer was done with the Blot[™] mini blot module (Invitrogen, Waltham, MA) following the manufacturer's protocol. Membrane was probed with mouse monoclonal anti-CRX antibody M02 (1:1 000, Abnova Corp., Taipei City, Taiwan) and rabbit polyclonal anti-HDAC1 antibody H51 (1:1 000, Santa Cruz Biotechnology, Dallas, TX), visualized with IRDye® 680RD goat anti-rabbit IgG and IRDye® 800CW goat anti-mouse IgG secondary antibodies (1:10 000, LI-COR, Inc., Lincoln, NE).

Transient transfection luciferase reporter assays

HEK293T cells were cultured in Dulbecco's Modified Eagle Medium supplemented with 10% fetal bovine serum (FBS) and Penicillin-Streptomycin following the manufacturer's protocol. Cells were transfected with calcium phosphate transfection protocol in 6-well plates as previously described^{17,18}. Experimental plasmids and usage amount are described in Supplementary Table S1. Typically, 48 hours after transfection, cells were harvested, digested and assayed for luciferase activity using Dual-Luciferase Reporter Assay System (Promega, Madison, WI) following the manufacturer's protocol. Data were collected using TD-20/20 Luminometer (Turner Designs, East Lyme, CT) and further processed with customized python scripts.

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Data Analysis

Homeodomain sequence alignment

The full-length protein sequences for the selected TFs were first aligned with Clustal Omega (EMBL-EBI, UK). Aligned sequences of the homeodomain helix three were then extracted to generate Figure 1B using Jalview (v2.11.1.7). A list of the accession numbers for the selected TFs can be found in Supplementary Table S1.

Determination of relative binding affinity with Spec-seq

For a biomolecular interaction between a protein P and a particular DNA sequence, S_i , the interaction can be diagrammed as:

$$P + S_i \rightleftharpoons P \cdot S_i \quad (1)$$

where $P \cdot S_i$ refers to the protein-DNA complex. The affinity of the protein *P* to sequence S_i is defined as the association constant K_A , or its reciprocal, the dissociation constant K_D . The K_A of the protein-DNA interaction is determined by measuring the equilibrium concentrations of each reactant and the complex:

$$K_A(S_i) = \frac{[P \cdot S_i]}{[P] \cdot [S_i]} \quad (2)$$

where [...] refers to concentrations. As in a typical Spec-seq experiment, thousands of different DNA sequences compete for the same pool of proteins, their relative binding affinities (the ratio of their K_A) can be determined by measuring the concentrations of each sequence in the bound and unbound fractions without measuring the free protein concentrations, which is often the most difficult to measure accurately:

$$K_A(S_1): K_A(S_2): \dots: K_A(S_n) = \frac{[P \cdot S_1]}{[S_1]}: \frac{[P \cdot S_2]}{[S_2]}: \dots: \frac{[P \cdot S_n]}{[S_n]}$$
(3)

In a binding reaction involving TF and a library of DNAs, the concentration of bound and unbound species are directly proportional to the number of individual DNA molecules in each fraction which can be obtained directly from sequencing data. With enough counts in each fraction, we can accurately estimate the ratios of concentrations from counts with the relationship:

$$\frac{[P \cdot S_i]}{[P \cdot S_j]} \approx \frac{N_B(S_i)}{N_B(S_j)} \text{ and } \frac{[S_i]}{[S_j]} \approx \frac{N_U(S_i)}{N_U(S_j)}$$
(4)

where N_U denotes counts in the unbound fraction and N_B denotes counts in the bound fraction. Therefore, the binding affinity of a sequence variant S_x relative to the reference sequence S_{ref} can be calculated by:

$$\frac{K_A(S_x)}{K_A(S_{ref})} \approx \frac{[P \cdot S_x]}{[P \cdot S_{ref}]} \frac{[S_{ref}]}{[S_x]} \approx \frac{N_B(S_x)}{N_B(S_{ref})} \frac{N_U(S_{ref})}{N_U(S_x)}$$
(4)

The natural logarithms of these ratios are the relative binding free energies in the units of *kcal/mol*. The relative free energy of the reference site for each CRX HD was set to 0.

Spec-seq data analysis and energy logo visualization

The sequencing results were first filtered and sorted based on conserved regions and barcodes. Reads with any mismatch in the conserved regions were discarded prior to further analysis as described previously^{32,86,87}. The ratio of individual sequence in bound and unbound reads was calculated as a measurement of relative binding affinity (Equation. 4) compared to the consensus sequence. The relative binding energy was then derived from the natural logarithm of the relative binding affinity and represented in *kcal/mol* units.

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For wild type CRX HD and all the mutants, the energy weight matrices (ePWMs) were generated based on the regression of the TF's binding energy to its reference sequence. Only sequences within two mismatches to the reference were used to generate the ePWMs. Energy logos were generated from ePWMs after normalizing the sum of energy on each position to 0 and the negative energy values were plotted such that preferred bases are on top. The sequence logos were generated from ePWMs with python package logomaker (v0.8). The ePWMs for all CRX HDs are listed in Supplementary Table S3.

ChIP-seq data analysis

2x150bp reads from Illumina NovaSeq were obtained for all samples with a minimum depth of 22M reads at Novogene (Beijing, China). For each sample, reads from two sequencing lanes were first concatenated and run through Trim Galore (v0.6.1)⁸⁸ to remove adapter sequences and then QC by FastQC (v0.11.5)⁸⁹. The trimmed reads were then mapped to the mm10 genome using Bowtie2 (v 2.3.4.1)⁹⁰ with parameters -X 2000 -very-sensitive. Only uniquely mapped and properly paired reads were retained with samtools (v1.9)⁹¹ with parameters -f 0x2 -q 30. Mitochondria reads were removed with samtools (v1.9)⁹¹. Duplicated reads were marked and removed with Picard (v2.21.4)⁹². Last, reads mapped to the mm10 blacklist regions were removed by bedtools (v2.27.1)⁹³ by intersect -v. bigWig files were generated with deeptools (v3.0.0)⁹⁴ and visualized on IGV Web App⁹⁵. For each genotype, an average binding intensity bigWig file from two replicates was generated with deeptools (v3.0.0)⁹⁴ command bamCompare –operation mean with default parameters.

Peak-calling was done with MACS2 (v2.1.1.20160309)⁹⁶ on individual replicate with the default parameters. For each genotype, we then generated a genotype specific high confidence peakset by intersection of peaks called in two replicates. IDR framework (v2.0.4)⁹⁷ were used to generate quality metrics for the processed ChIPseq data. R package DiffBind (v3.0.15)^{98,99} and DEseq2 (v1.30.1)¹⁰⁰ were then used to re-center peaks to ± 200 bp regions surrounding summit, generate normalized binding intensity matrix, and differential binding matrix. We defined differentially bound peaks between each mutant and wild type sample if the absolute log₂FC is more than 1.0, corresponding to two-fold, and the FDR is smaller than 5e-2.

To associate peaks to genes, we used Genomic Regions Enrichment of Annotations Tool (GREAT v4.0.4)¹⁰¹ through the R package rGREAT (v1.19.2)¹⁰². Each peak was assigned to the closest TSS within 100kb.

Binding intensity heatmap and clustering

To generate the binding intensity heatmap in Figure 2A, we first compiled the genotype specific high confidence peakset for all genotypes into a single consensus peakset and only peaks with at least 5 cpm in all genotypes were retained. Python package fastcluster $(v1.1.26)^{103}$ was used to perform hierarchical clustering of the consensus peakset intensity matrix with parameters method='single', metric='euclidean'. The genomic regions corresponding to the two major clusters were exported and used to generate binding intensity heatmaps with deeptools $(v3.0.0)^{94}$.

Genomic region enrichment of CRX peaks

Peak annotation in Figure 2D were obtained using annotatePeaks.pl from HOMER (v4.8).

De novo motif searching

The mm10 fasta sequences for each genotype specific peaks were obtained using R package BSgenome (v 1.58.0)¹⁰⁴. De novo motif enrichment analysis for each set of sequences were then performed with MEME-ChIP in MEME Suite (v5.0.4)¹⁰⁵ using order 1 Markov background model and default parameters. Since homeodomain motifs are relatively short and can be repetitive (e.g. K88N motif), we reported DREME³⁷ found motifs for Figure 2E, which is more sensitive than MEME to find short, repetitive motifs.

<u>RNA-seq data analysis</u>

2x150bp reads from Illumina NovaSeq were obtained for all samples with a minimum depth of 17 M reads at Novogene (Beijing, China). Sequencing reads were first run through Trim Galore (v0.6.1)⁸⁸ to remove adapter

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sequences and then QC by FastQC $(v0.11.5)^{89}$. Trimmed reads were then mapped to the mm10 genome and quantified with kallisto $(v0.46.2)^{106}$. Kallisto output transcript-level abundance matrices were then imported and summarized into gene-level matrices with R package tximport $(v1.18.0)^{107}$. DEseq2 $(v1.30.1)^{100}$ was then used for normalization and differential expression analysis. The normalized count and differential expression matrices were then exported and further processed with customized python scripts.

We defined differentially expressed genes between each mutant and wild type sample if the absolute log_2FC is more than 1.0, corresponding to two-fold, and the FDR is smaller than 1e-2. For comparison between heterozygous and homozygous mutants, we first filtered genes with at least 5 cpm and then those that were called differentially expressed compared with wild type in at least one mutant genotype. We retrieved gene names in supplementary tables 3-5 from the Database for Annotation, Visualization and Integrated Discovery (DAVID, v6.8)¹⁰⁸.

Definition of CRX-dependent and -independent gene set

We first identified CRX peaks that were bound in the *WT* sample but lost in *R90W/W* sample (log₂FC < -1 and FDR < 5e-2). This yielded a total of 7677 peaks. We then found the genes associated with these peaks. We defined a gene to be CRX-dependent activated if its expression was down in adult (P21) *R90W/W* RNA-seq sample (log₂FC < -0.6 and FDR < 1e-5). Similarly, a gene is defined as CRX-dependent suppressed if its expression was up in adult *R90W/W* RNA-seq sample (log₂FC > 0.6 and FDR < 1e-5). A gene is defined as CRX-independent if its expression was not significantly affected in adult *R90W/W* RNA-seq samples. There were 617 CRX-dependent activated, 135 CRX-dependent suppressed, and 5565 CRX-independent genes. Manual inspection of the CRX-dependent suppressed genes revealed no clear association with photoreceptor development. Therefore, we did not further pursue this gene set. The complete list of CRX-dependent activated genes that showed differential expression in at least one of the HD mutant retinas can be found in Supplementary Table S5. The lists for CRX-independent genes that showed differential expression in Supplementary Tables S6 and S7 respectively.

Gene ontology analysis

Gene ontology analysis in Figures S4B, S4E, and S5F were performed using R package clusterProfiler $(v4.0.5)^{109,110}$ with the genome wide annotation package org.Mm.eg.db $(v3.12.0)^{111}$. Redundant enriched GO terms were removed using simplify() function with parameters cutoff=0.7, by="p.adjust". The enrichment analysis results were then exported in table format and further processed for plotting with python.

Aldiri et al. RNA-seq data re- analysis

The RNA-seq data from *Aldiri* et al.³⁸ were obtained from GEO under accession numbers GSE87064. The reads were processed similarly as all other RNA-seq data generated in this study. For Figures S4C, S4F, S5C, S5E,S6A, S6B, expression row z-scores were calculated using average cpm from replicates at each age.

Statistical analysis

One-way ANOVA with Turkey honestly significant difference (HSD) test in Figures S3C and 6A were performed with python packages scipy $(v1.8.1)^{112}$ and scikit_posthocs $(v0.7.0)^{113}$. Two-sided Mann-Whitney U test in Figure S5A was performed with python package scipy $(v1.8.1)^{112}$.

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Figure 1. Disease associated missense mutations altered CRX HD DNA binding specificity. (A) Diagram of CRX functional domains: Homeodomain (HD) for DNA-binding and Activation Domain (AD) for target gene transactivation. The three missense mutations in this study are located at the C-terminus of CRX HD and associated with different retinal diseases in human. Number in the parenthesis denotes the CRX HD position of the corresponding mutated residue. (B) Alignments of HD recognition helix sequences for the indicated HD proteins for which HD missense mutations have been associated with inherited diseases. Accession numbers can be found in Supplementary Table S1. Missense variants in this study (highlighted) are located at highly conserved residues across species and different HD TFs. (C) Spec-seq experimental workflow (STAR Methods). (D) Spec-seq library design of monomeric HD binding sites. (E) EMSA gel images of Spec-seq experiments with different CRX HD species. B_x: Bound. B.: Unbound. (F) Relative binding energy comparison from two different experiments with WT HD. (G) Binding energy model for WT CRX HD. (H-J) Relative binding energy comparison between WT HD and R90W HD (H), E80A HD (I), or K88N HD (J). Consensus sequence is defined to have relative binding energy of 0kT (TAATCC for WT, R90W and E80A, TAATTA for K88N). The diagonal line of identity is represented in grey dash. The orange dashed line shows the best linear fit to the data. (K-M) Binding energy models for R90W HD (K), E80A HD (L), and K88N HD (M). Only sequence variants within two mismatches to the corresponding consensus sequences were used to generate binding models. Negative binding energy is plotted such that bases above the x-axis are preferred bases and bases below the x-axis are unfavorable bases. Constant bases (TAA) carried no information are drawn at arbitrary height in grey.



Figure 2. CRX E80A binds to WT sites while CRX K88N occupies novel genomic regions enriched for N88 HD motif *in vivo.* (A) Enrichment heat map depicting CRX ChIP-seq normalized reads centered at all possible CRX peaks ± 2kb, sorted by binding intensity in *WT* samples. Clusters were defined by hierarchical clustering of CRX binding intensity matrix from all genotypes (STAR Methods). (B-C) Genome browser representations of chIP-seq normalized reads for different CRX species in P14 *WT* and mutant mouse retinas at *Rho* and *Atf2*. (D) Enrichment heatmap showing fraction of CRX ChIP-seq peaks fall in different genomic environments. (E) Logo representations of *de novo* found short HD motifs under CRX ChIP-seq peaks in *WT* and mutant mouse retinas with DREME *E-value* on the right.



Figure 3. CRX-dependent activated genes affected in opposite directions in developing *Crx^{E80A}* and *Crx^{K80N}* mutant retinas (A) Heat map showing sample-wise Pearson correlations of the expression of all CRX-dependent activated genes between P10 *WT* and HD mutant mouse retinas in this study (rows) with post-natal *WT* retinas from age P3 to P21 (columns, data from GSE87064). (B) Heat map showing the expression changes of DEGs in CRX-dependent activated gene set in HD mutant mouse retinas at P10. (C-D) Heat maps showing expression changes of selected photoreceptor genes from Group 1 and Group 2. Color scale identical to (B).



Figure 4. Photoreceptor genes important for phototransduction are down-regulated in all HD mutants (A) Box plot showing that genes in the detection of light stimulus GO term were down-regulated and affected to various degrees in different adult (P21) HD mutant mouse retinas. (B) Heat map showing that expression of both cone and rod phototransduction genes were down-regulated in adult (P21) HD mutant mouse retinas. Annotation of rod and cone enrichment of each gene is in Supplementary Table 6. See Supplementary Figure S6 for the developmental expression dynamics of these genes.



Figure 5. Only *Crx^{E80/4/*}* retinas maintain photoreceptor OS and residual rod ERG response (A-E) Hematoxylin-eosin (H&E) staining of P21 retina sections show that photoreceptor OS layer is absent in all mutant retinas except *Crx^{E80/4/*}*. OS: outer segment; ONL: outer nuclear layer; INL: inner nuclear layer; IPL: inner plexiform layer; GCL: ganglion cell layer. Scale bar, 100µm. (F-J) Rhodopsin (RHO, red) immunostaining is present in *Crx^{E80/4/*}*, *Crx^{E80/4/*}*, and *Crx^{K80/4/*}* retinas and absent in *Crx^{K80/V*}* retina. Cone arrestin (mCAR, green) immunostaining is absent in all mutant retinas. Nuclei were visualized by DAPI staining (Blue). Scale bar, 100µm. (K-M) The electroretinogram responses (ERG) recorded from 1-month mice. Error bars represent the standard error of the mean (SEM, n ≥ 4). *p*-value: Two-way ANOVA and Tukey's multiple comparisons. ****: p ≤ 0.0001. ns: >0.05.



Figure 6. CRX E80A hyperactivity underlies precocious photoreceptor differentiation in Crx^{E80A} retinas. (A) Boxplot showing luciferase reporter activities of different CRX variants. *P-values* for one-way ANOVA with Turkey honestly significant difference (HSD) test are indicated. *p-value*: ****: ≤0.0001, ***: ≤0.001, ns: >0.05. (B-D) Rhodopsin (RHO, green) immunostaining is absent in P3 *WT* retina but detected in $Crx^{E80A/4}$ and $Crx^{E80A/4}$ retinas. Nuclei are visualized by DAPI staining (Blue). Arrow indicates the sporadic RHO staining in $Crx^{E80A/4}$ sample. ONBL: outer neuroblast layer; GCL: ganglion cell layer. Scale bar, 100µm.



Figure 7. Missense mutations in CRX HD affect photoreceptor gene expression and leads to distinct retinal disease phenotypes through gain- and loss-of-function mechanisms.



Supplementary Figure S1. Multi-omics approach to investigate the functional consequences of dominant disease mutations on CRX regulatory activities and photoreceptor development. Human retinopathy associated CRX HD mutant is first tested *in vitro* for HD-DNA interactions by Spec-seq. Quantitative binding models

Human retinopathy associated CRX HD mutant is first tested *in vitro* for HD-DNA interactions by Spec-seq. Quantitative binding models are generated for WT and mutant HDs. Each mutation is then introduced into endogenous *mCrx* locus to generate human mutation knock-in mouse models. ChIP-seq is employed to characterize CRX chromatin binding in *WT* and mutant mouse retinas. Bulk RNA-seq is then applied to determine transcriptomic changes in developing (P10) and mature (P21) photoreceptors in both *WT* and mutant mouse retinas. Last, phenotypic characterization on retinal morphology and visual functions is carried out to understand the consequences of mutant CRX chromatin binding and associated transcriptomic alterations.



Supplementary Figure S2. Reversed-strand Spec-seq library showed similar changes in mutant CRX HD DNA-binding specificity. (A) Native SDS-PAGE gel image of affinity purified empty GST tag and GST-CRX HDs. (B-D) Relative binding energy comparison from two different experiments for R90W HD (B), E80A HD (C), and K88N HD (D) on the same

(B-D) Relative binding energy comparison from two different experiments for R90W HD (B), E80A HD (C), and K88N HD (D) on the same Spec-seq library as in Figure 2.

(E-L) Spec-seq experiments of a second library with the TAANNN sites on the reverse strand show similar results. (E-H) Relative binding energy comparison from two different experiments for WT HD (E), R90W HD (F), E80A HD (G), and K88N HD (H) on the reversed monomeric library. The diagonal line of identity is represented in grey dash. The orange dashed line shows the best linear fit to the data. (I-L) Binding energy models for WT HD (I), R90W HD (J), E80A HD (K), and K88N HD (L) obtained from the reversed monomeric library. Quantitative difference with models obtained from forward-oriented library likely comes from difference in sequences immediately flanking the TAANNN variable region.



Supplementary Figure S3. WT and mutation knock-in mouse CRX sequences and genotyping identifications

(A) Alignment of *mCrx* cDNA and protein sequences showing the nucleotide substitutions and amino acid changes of *Crx^{E80A}* (top) and *Crx^{K88N}* (bottom) alleles. Only coding regions of *mCrx* exons are shown and the diagram is not to scale. Underlined bases in WT sequences indicate the restriction enzyme Hinfl cut sites used in the genotyping PCR.

(B) Representative mutation knock-in mouse genotyping gel image.

(C) Barchart and stripplot showing mCrx mRNA expression levels in P14 WT and mutant mouse retinas. P-values for one-way ANOVA with Turkey honestly significant difference (HSD) test are indicated. p-value: ****: ≤0.0001, ***: ≤0.001, **: ≤0.01, *: ≤0.05, ns: >0.05.
(D) Immunoblots of nuclear extracts generated from P14 WT and mutant mouse retinas showing that full-length CRX protein are produced and localized to the nucleus fraction in all mutant mouse retinas. HDAC1 was used as a loading control.



Supplementary Figure S4. Definition of CRX-dependent activated and CRX-independent gene sets.

(A) Schematic representation of CRX-dependent activated genes where CRX binding nearby is required for the expression of these genes in mature WT retinas.

(B) Top GO terms associated with CRX-dependent activated genes. Benjamini-Hochberg adjusted p-values are shown. (C) Line plot showing average expression pattern of CRX-dependent activated genes during normal post-natal retina.

(D) Definition of CRX-independent genes where CRX binding nearby is dispensable for the expression of these genes in mature WT retinas.

(E) Top GO terms associated with CRX-independent genes. (F) Line plot showing average expression pattern of CRX-independent genes during normal retina development. RNA-seq data in (C) and (F) were retrieved from GSE87064.



Supplementary Figure S5. E80A and K88N mutation each causes novel gene expression changes in the CRX-independent category (A) Strip plots showing normalized CRX ChIP-seq intensity at peaks associated with Group1 or Group2 genes in WT and CRX mutant mouse retinas. *P-values* for two-sided Mann-Whitney U test are indicated. *p-value*: ****: ≤ 0.001 , **: ≤ 0.001 , **: ≤ 0.001 , **: ≤ 0.005 , ns: > 0.05. (B) Venn diagram showing the overlap of genes differentially expressed (DEGs) in Crx^{E80A} (pale blue) and Crx^{KBBN} (pale yellow) but not in Crx^{F90WW} (grey) mutant retinas. For DEGs in Crx^{E80A} and Crx^{KBBN} mutants, genes that were differentially expressed in either heterozygotes or homozygotes, or both were counted.

(C) Heat map showing the expression changes of CRX-independent genes that are DEGs in at least one of the Crx^{E80A} mutants (n = 244, left). Heat map on the right shows the expression pattern of these genes during normal post-natal development (data from GSE87064).
(D) Table showing selected genes down-regulated in Crx^{E80A} mutants that have been implicated in cell differentiation or photoreceptor development.

(E) Heat map showing the expression changes of CRX-independent genes that are DEGs in at least one of the *Crx^{K88N}* mutants (n = 351, left). Heat map on the right shows the expression pattern of these genes during normal post-natal development (data from GSE87064). (F) Bar chart showing GO term enrichment of DEGs in *Crx^{K88N}* mutants.



Supplementary Figure S6. Developmental expression pattern of phototransduction genes in WT animals

(A) Parallel coordinates plot showing expression pattern of genes in GO:0009583 during normal post-natal retina development (data from GSE87064). Normalized row z-score is shown

(B) Heatmap showing expression patterns of phototransduction genes during normal post-natal retina development (data from GSE87064). Normalized row z-score is shown.