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2017

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### Recommended Citation

Ruzyski, Philip A.; Linne, Courtney D.; Hennig, Anne K.; and Chen, Shiming, "Crx-L253X mutation produces dominant photoreceptor defects in TVRM65 mice." *Investigative Ophthalmology & Visual Science*. 58, 11. 4644–4653. (2017).

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# Crx-L253X Mutation Produces Dominant Photoreceptor Defects in TVRM65 Mice

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Submitted: April 19, 2017

Accepted: August 3, 2017

Citation: Ruzycski PA, Linne CD, Hennig AK, Chen S. *Crx-L253X* mutation produces dominant photoreceptor defects in *TVRM65* mice. *Invest Ophthalmol Vis Sci*. 2017;58:4644-4653. DOI:10.1167/iovs.17-22075

**PURPOSE.** The cone-rod homeobox (CRX) transcription factor is essential for photoreceptor gene expression, differentiation, and survival. Human *CRX* mutations can cause dominant retinopathies of varying onset and phenotype severity. In animal models, dominant frameshift *Crx* mutations introduce a premature termination codon (PTC), producing inactive truncated proteins that interfere with normal CRX function. Previously, a mutant mouse, *TVRM65*, was reported to carry a recessive late PTC mutation, *Crx-L253X*. More detailed phenotype analysis of the pathogenicity of *Crx-L253X* sheds new light on the variability of *CRX*-linked diseases.

**METHODS.** Homozygous (*L253X/X*); heterozygous (*L253X/+*); *Crx*<sup>-/-</sup> and control *C57BL/6J* (WT) mice were analyzed at various ages for changes in retinal function (ERG), morphology (histology) and photoreceptor gene expression (qRT-PCR).

**RESULTS.** At 1 month, *L253X/X* mice lack visual function, show greater reductions in retinal thickness, and distinct gene expression changes relative to *Crx*<sup>-/-</sup>, suggesting that the phenotype of *L253X/X* is more severe than *Crx*<sup>-/-</sup>. *L253X/+* mice have reduced rod/cone function, but normal retinal morphology at all ages tested. qRT-PCR assays described a complex phenotype in which both developing and mature photoreceptors are unable to maintain proper gene expression. *L253X* mRNA/protein is overexpressed relative to normal *Crx*, suggesting a pathogenic mechanism similar to early PTC mutations. However, the overexpression is less pronounced, correlating with a relatively mild dominant phenotype.

**CONCLUSIONS.** The *L253X* mouse provides a valuable model for *CRX*-associated retinopathy. The pathogenicity of *CRX* frameshift mutations depends on the position of the PTC, which in turn determines the degree of mutant mRNA/protein overproduction.

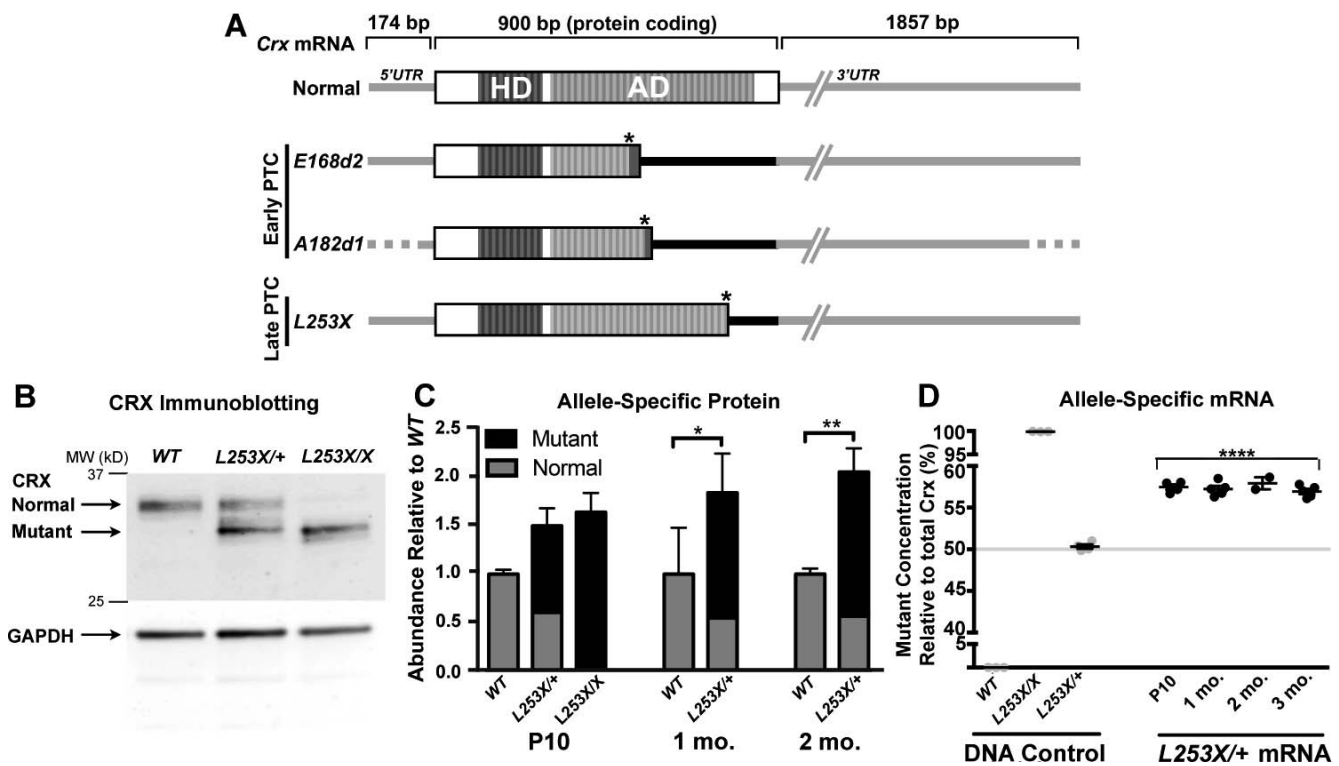
**Keywords:** *CRX* mutations, animal models, inherited photoreceptor degeneration, cone-rod dystrophy, gene expression

The cone-rod homeobox protein (CRX) is a transcription factor (TF) that regulates expression of many photoreceptor genes essential for the development and maintenance of both rod and cone photoreceptors.<sup>1-4</sup> Human *CRX* mutations are associated with various diseases, including retinitis pigmentosa (RP); cone-rod dystrophy (CoRD); and Leber congenital amaurosis (LCA).<sup>5,6</sup> *CRX*-linked diseases are largely inherited in an autosomal dominant fashion or arise de novo, and vary widely in severity and age of onset.<sup>5,7-10</sup> Since *CRX* mutations can be detected with early genetic testing, we need to be able to predict their effects and to define effective treatment and gene therapy regimens.<sup>7</sup>

Previous research has divided disease causing *CRX* mutations into four classes.<sup>11</sup> Classes I and II are missense mutations that fall within or near the region coding for the DNA-binding homeodomain; class I mutations reduce the binding of CRX to its DNA targets.<sup>12</sup> Previous work has shown by a variety of metrics that heterozygous mice carrying the class I *R90W* mutation suffer from a mild form of CoRD, similar to human patients with such mutations, while homozygotes exhibit an LCA-like condition.<sup>13</sup> Class III and IV mutations represent frameshift mutations caused by insertions or deletions in the region coding for the

transactivation domains of CRX.<sup>12</sup> Class IV mutations, modeled by the *RIP* mouse, cause translation of a much longer peptide sequence due to a frameshift and extension of the open reading frame (ORF) into the 3' untranslated region (3'UTR).<sup>14</sup> In contrast to this extended peptide sequence, Class III mutations truncate the ORF of *CRX* with a premature termination codon (PTC) that results in a CRX protein with a shortened transactivation domain and leads to a severe dominant degenerative phenotype (LCA or CoRD).<sup>13</sup> This phenotype has been modeled by the *E168d2* mouse<sup>13</sup> and *Rdy* (*A182d1*) cat<sup>15</sup> (Fig. 1A). In vitro DNA binding and transactivation experiments determined that the *E168d2* protein has a similar affinity for CRX target DNA sequences as normal CRX, but is unable to activate transcription on its own or in combination with other retinal TFs that normally synergize with CRX.<sup>12,13</sup> Furthermore, when tested together with normal CRX, *E168d2* protein interfered with normal CRX transactivation in a dominant-negative manner.<sup>13</sup>

Class III PTC-causing *CRX* mutations also result in a novel untranslated region between the PTC and normal stop codon, which becomes a part of the mutant mRNA transcript's 3'UTR (Fig. 1A).<sup>11</sup> In the *E168d2* mouse and the *Rdy* (*A182d1*) cat, longitudinal studies showed gradual accumulation of mutant



**FIGURE 1.** Class III *Crx* mutations introduce a PTC, resulting in accumulation of truncated CRX proteins and mutant mRNA with variable extended 3'UTR. (A) Schematic representation of normal and mutant *Crx* mRNA species, showing untranslated (UTR, gray/black lines) and protein-coding (open/stripped boxes) with their relative sizes (bp), based on mouse (Normal, E168d2<sup>13</sup> and L253X) and cat (A182d1)<sup>15</sup> models. The DNA-binding homeodomain (HD, dark strips) and the activation domain (AD, light strips) sequences<sup>12</sup> are indicated. All three mutant mRNAs encode the complete HD, but ADs are truncated at different positions due to their PTC (indicated as the right-hand end of the protein coding box) induced by the mutation (marked by an asterisk). These PTCs also expand the 3'UTR to include different lengths of the original coding region (indicated by black line) in front of the normal 3'UTR (gray line). The position of the PTC relative to the normal stop codon determines the length of expanded 3'UTR: Early PTC mutants, E168d2 and A182d1 (*Rdy* cat mutation) produce a longer 3'UTR than the late PTC mutant L253X. Since 5'UTR and 3'UTR sequences of feline *Crx* mRNA are not available, these regions of the A182d1 transcript are indicated by dotted lines. (B) Immunoblot of CRX in the retinas of WT, L253X/+, and L253X/X mutants at 1-month old (mo) with GAPDH as a protein loading control. Black arrows indicate the running position for the normal and mutant (truncated) forms of CRX. (C) Quantification of relative amounts of normal and mutant CRX proteins in the indicated retinas at three ages (mean  $\pm$  SEM,  $n \geq 3$ ; \*  $P < 0.05$ , \*\*  $P < 0.01$ ; 2-Way ANOVA with Tukey's multiple comparisons test). (D) ddPCR quantification of mutant (L253X) and normal *Crx* mRNA as percentage of total *Crx*. The specificity of the two allele-specific ddPCR assays was established using triplicate tail DNA samples (gray circles) from the indicated genotypes (DNA control). L253X/+ mRNA results are presented as percent of total *Crx* mRNA (mutant plus normal transcripts; mean  $\pm$  SEM,  $n \geq 3$ ; \*\*\*\*  $P < 0.0001$ ; unpaired *t*-test for mutant mRNA level relative to L253X/+ DNA control;  $n \geq 3$ ).

protein and mRNA.<sup>13,15</sup> The mechanism for this accumulation is unknown, but was postulated to arise from decreased mutant RNA degradation,<sup>13</sup> possibly due to the presence of cryptic stabilizing elements within the PTC-expanded mutant 3'UTR. Therefore, the position of the PTC (relative to the normal termination codon) could determine the stability, and thus abundance of mutant RNA.

Previously, another PTC mutant mouse, *TVRM65* (*Crx*:L253X), was discovered in a chemical-induced mutagenesis screen conducted by The Jackson Laboratory<sup>16</sup> (Bar Harbor, ME, USA; see Methods section for nomenclature specifics). The L253X mutation (Fig. 1A) occurs later (closer to the normal termination codon) than E168d2 and A182d1. L253X would be a candidate to model human late PTC class III mutations, such as L237I<sup>17</sup> and Q256X<sup>18</sup> associated with dominant CoRD. However, the initial histologic assessments suggested that the L253X mouse phenocopies the recessive null (*Crx*<sup>-/-</sup>) condition.<sup>16</sup> Here, we describe more detailed phenotypic analysis of L253X mice, which demonstrates that this mouse models a new dominant but mild form of class III disease. Our results support the hypothesis that class III mutation pathogenicity depends on the position of the mutation-induced PTC (a late PTC produces a less severe phenotype than an early PTC),

which correlates with the abundance of mutant protein products. These findings have implications for predicting human *CRX* disease severity and progression.

## METHODS

### Nomenclature for the *TVRM65* Mutation

There are several annotated transcripts for the murine *Crx* locus. The previously published amino acid change in the *TVRM65* mouse was based on isoform 2 (NM001113330.1) that encodes a CRX protein with 323 amino acids.<sup>16,19</sup> However, the major transcript in the retina is isoform 1 (NM\_007770.4), which encodes a CRX protein homologous to human CRX, with 299 amino acids. Thus, here we refer to the *TVRM65* mutation as *p.L253X*, based on isoform 1 numbering.

### Mice

*Crx*-L253X (*TVRM65*) mice<sup>16</sup> were obtained from The Jackson Laboratory (MGI ID: 4867395). Mice were mated and maintained with C57BL/6J (JAX Stock number 000664), and

confirmed free of *rd1* and *rd8* mutations by PCR genotyping. *Crx* null mice (*Crx*<sup>-/-</sup>) were provided by Constance Cepko, Harvard University (Boston, MA, USA)<sup>20</sup> and back-crossed onto *C57BL/6J* for more than 10 generations. All mice were housed in a barrier facility in the Division of Comparative Medicine of Washington University School of Medicine. All procedures involving the use of mice were approved by Washington University's Animal Care and Use Committee and followed the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

PCR genotyping, RNA purification and reverse transcription, and qRT-PCR were performed as previously described.<sup>13</sup> Primer sets for genotyping and RT-PCR are listed in Supplementary Table S1. For qRT-PCR, all primers were tested for proper amplification efficiency prior to use. Relative gene expression was normalized to the retinal constitutively expressed genes *Ubb*, *Tuba1B*, and *Gapdh*. Data for 3 biologic replicates were then analyzed using the Delta Cq method in qPCR analysis software (QBase; Biogazelle, Ghent, Belgium). The results are presented by the heatmap.2 function of the R gplots package (v3.0.1).

### Droplet Digital PCR (ddPCR)

Confirmatory genotyping and allelic expression data were generated using the droplet digital PCR system (BioRad, Hercules, CA, USA). A 20  $\mu$ L ddPCR reaction mixture containing custom-designed normal and mutant primer/probe sets (see Supplementary Table S2) was prepared according to manufacturer's directions, droplets generated, and nanoreactions cycled on a thermal cycler (C1000 Touch; Bio-Rad Laboratories). The average number of allelic transcripts per biological replicate ( $n = 3$ ) was then determined with a droplet reader and analytical software (QX 200 with QuantaSoft Analysis Pro; Bio-Rad Laboratories).

### Transient Transfection Luciferase Reporter Assays

HEK293T cells (ATCC CRL-11286) were cultured in Dulbecco's modified Eagle medium with 10% fetal bovine serum and penicillin/streptomycin. Cells were transfected using a conventional CaCl and boric acid buffered saline method in 6-well plates as previously described.<sup>12,13</sup> Experimental plasmids and transfection amounts included 2  $\mu$ g *BR130-Luc* (a *Rbo* promoter-luciferase reporter)<sup>12</sup>; 1 ng *pRL-CMV* (transfection normalization control); and protein expression vectors *pED-NRL* (100 ng), *pcDNA3.1-bCRX*, *CRX1-254*, and *CRX-E168d2*. Cells were harvested 48 hours posttransfection and assays performed using the dual-luciferase reporter assay system (Promega Corp., Madison, WI, USA).

### ERG and Statistical Analyses

These were performed as previously described.<sup>13</sup> Briefly, tests were performed on a visual electrodiagnostic system (UTAS-E3000 with EM for Windows; LKC Technologies, Inc., Gaithersburg, MD, USA) while mouse body temperature was maintained at 37°C  $\pm$  0.5°C with a heating pad controlled by a rectal temperature probe (FHC, Inc., Bowdoin, ME, USA). Pupils were dilated with 1.0% atropine sulfate (Bausch & Lomb, Tampa, FL, USA) and dilation and corneal hydration maintained during testing by positioning the platinum wire loop recording electrodes in a mixture of atropine and 1.25% hydroxypropyl methylcellulose (GONAK; Akorn, Inc., Buffalo Grove, IL, USA). Mice were tested without knowledge of genotype. Bilateral flash ERG responses were obtained; the set of recordings giving the higher amplitudes was correlated with genotype information for statistical analyses including 2-way ANOVA and post-hoc

multiple comparison tests were performed using graphing software (GraphPad Prism; GraphPad Software, Inc., La Jolla, CA, USA).

### Quantitative Western Blot Analysis

As previously described,<sup>13</sup> experiments were performed using three biologic replicates (15  $\mu$ g total protein extract from homogenates of two retinas). Membranes were probed with mouse monoclonal anti-CRX antibody M02 (1:200; Abnova Corp., Taipei City, Taiwan) and anti-GAPDH antibody (1:10,000, G9545; Sigma-Aldrich Corp., St. Louis, MO, USA), visualized with donkey anti-rabbit 680 nm and donkey anti-mouse 800 nm fluorescent dye (1:10,000 IRDyes; LI-COR, Inc., Lincoln, NE, USA), then imaged and quantified using commercial software (Odyssey Infrared Imager and ImageStudio 6; LI-COR, Inc.).

### Histology, Morphometry, Immunohistochemistry, And Statistical Analyses

These assays were performed as previously described.<sup>13</sup> Sagittal sections (4- $\mu$ m) of paraffin-embedded eyecups, cut through the optic nerve head, were either stained with hematoxylin and eosin (H&E) or prepared for immunofluorescence staining. Slides containing sections were incubated overnight with primary antibody at 4°C (1:400 anti-Rhodopsin monoclonal RetP1; Sigma-Aldrich Corp.), washed, and incubated with secondary antibody (1:500 AlexaFluor 568 goat-anti-mouse; Invitrogen Corp., Carlsbad, CA, USA) for 2 hours at room temperature. Coverslips were mounted with mounting medium with DAPI (Vectashield Hardset; Vector Laboratories, Inc., Burlingame, CA, USA). All imaging was performed using a microscope and camera (DM5500B and DFC365FX; Leica Microsystems GmbH, Buffalo Grove, IL, USA).

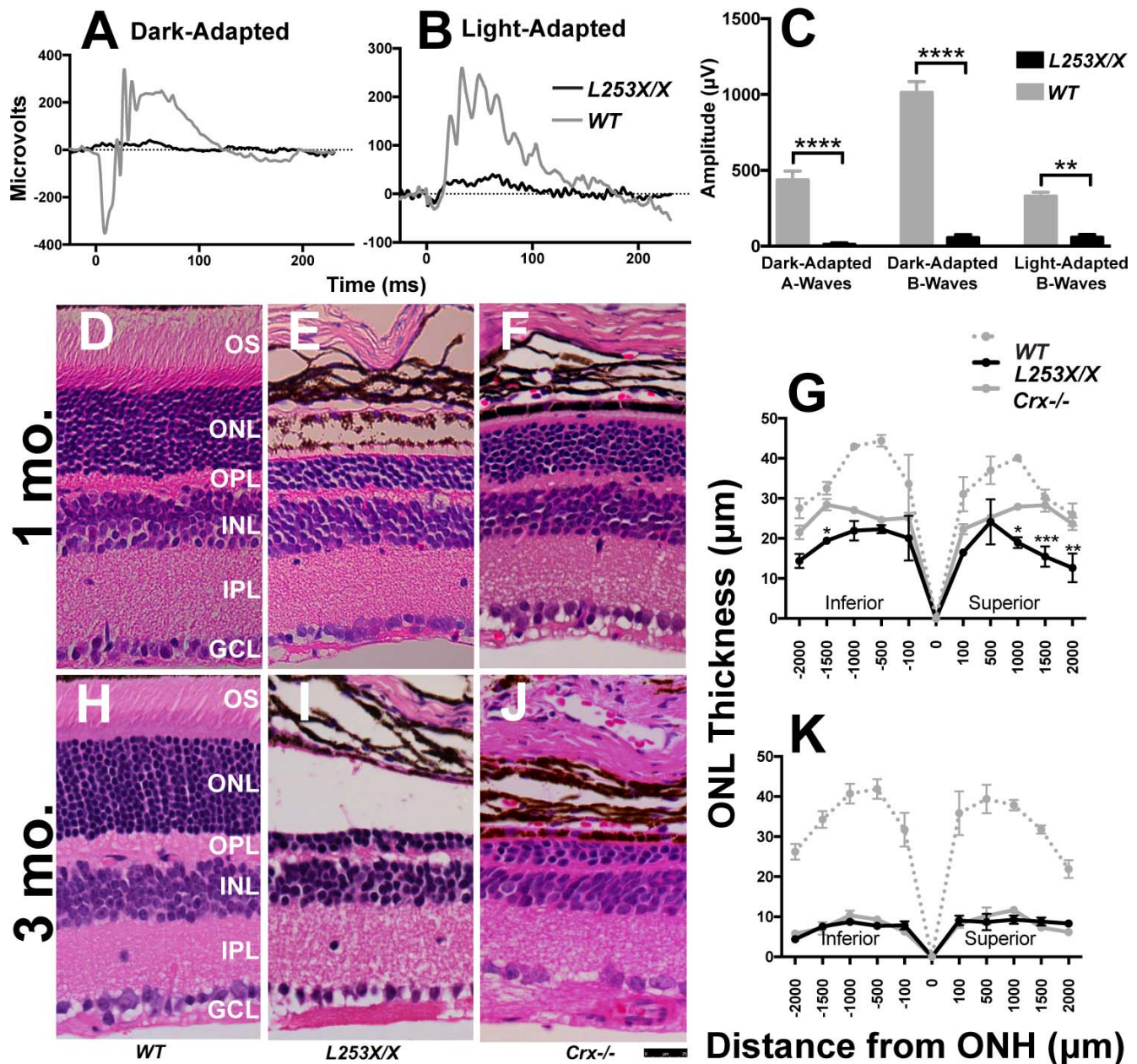
## RESULTS

### L253X Overproduces Mutant mRNA/Protein in Affected Retinas

A hallmark of class III *CRX* frameshift mutations is the overexpression of the mutant allele.<sup>13,15</sup> To determine if *L253X* retinas accumulate mutant protein, we performed quantitative Western blots on total protein extracts from retinas of *WT* and heterozygous *L253X/+* mice at postnatal day 10 (P10), 1 and 2 months old, and homozygous *L253X/X* mice at P10 only (because photoreceptors degenerate before later ages).<sup>16</sup> Immunoblotting with an anti-CRX antibody revealed a truncated *L253X* protein that ran faster than the normal full-length *CRX* protein in samples from both heterozygous and homozygous mutant retinas (Fig. 1B). We quantified the total amount of *CRX* protein and of each isoform. Both *L253X/+* and *L253X/X* displayed a 1.5- to 2.2-fold increase in total *CRX* protein compared to the *WT* retinas (Fig. 1C). The increases can be attributed to an accumulation of mutant protein (black bar), since normal *CRX* protein levels (gray bar) in *L253X/+* retinas with one normal allele were roughly half of the total *CRX* protein present in *WT* retinas with two alleles.

To determine if the increased mutant protein corresponded with an increase in mutant mRNA, we quantified the relative allelic expression of *L253X* and normal *Crx* mRNA using droplet digital PCR (ddPCR). First, using mutant and *WT* mouse DNA, we developed a ddPCR assay that clearly distinguished the *L253X* and normal *Crx* sequences (Fig. 1D, DNA control). This assay was then used to quantify allele-specific mRNA species. At all ages tested, the mutant transcript was





**FIGURE 2.** *L253X/X* retinas lack rod and cone responses to light and degenerate earlier than *Crx*<sup>-/-</sup>. ERG was used to measure rod (dark-adapted) and cone (light-adapted) responses to various light stimuli in 1-mo *L253X/X* and WT mice. (A, B) Comparison of representative ERG traces at the highest light intensity: 0.89 log scot cd/m<sup>2</sup> for dark-adapted responses, and 2.67 log scot cd/m<sup>2</sup> for light-adapted responses. (C) Bar graphs compare mean A-wave and/or B-wave amplitude differences between the two genotypes at the highest light intensity tested (mean ± SEM, *n* ≥ 3; \* *P* ≤ 0.05, \*\*\* *P* ≤ 0.0005, \*\*\*\* *P* < 0.0001; 2-way ANOVA with Sidak's multiple comparisons test). (D–F, H–J) H&E-stained paraffin-embedded sagittal retinal sections from the indicated mice at 1 and 3 mo. Images were taken approximately 500 μm from the ONH at a ×40 magnification (scale bar: 25 μm). (G, K) Morphometry measures at five set distances (in microns) on either side of the ONH (mean ± SEM, *n* ≥ 3; \* *P* ≤ 0.05, \*\* *P* < 0.01, \*\*\* *P* ≤ 0.0003; 2-way ANOVA with Sidak's multiple comparisons test).

overrepresented, accounting for 57% of total *Crx* mRNA (Fig. 1D). Total *Crx* transcript levels quantified by conventional qRT-PCR (Supplementary Fig. S1) showed a statistically significant total *Crx* mRNA increase in 1-month *L253X/+* retinas relative to WT, although levels were comparable to WT retinas during development at P10 and by 3 months.

Overall, *L253X/+* retinas develop and maintain an elevated level of mutant mRNA that translates into an imbalance of the two protein forms, but this misregulation is not as severe as observed in other class III mutants. These results identify *Crx*-

*L253X* as a bona fide class III mutation, but suggest that the position of the PTC determines the toxicity of the allele.

### *L253X/X* Mice Have No Detectable Rod and Cone Function

Homozygous animals carrying class III or null *Crx* mutations lack measurable photoreceptor function.<sup>13,15,20</sup> To confirm that *L253X/X* photoreceptors are functionally compromised, we quantified rod/cone light responses in *L253X/X* and WT control mice at 1 month using ERG. Even at the highest light intensity stimuli, *L253X/X* retinas displayed very little

response (Figs. 2A–C). These results confirm that, like other class III models, homozygous *L253X/X* animals lack photoreceptor visual responses, resembling human LCA.

### *L253X/X* Photoreceptors Degenerate Earlier Than *Crx*<sup>−/−</sup>

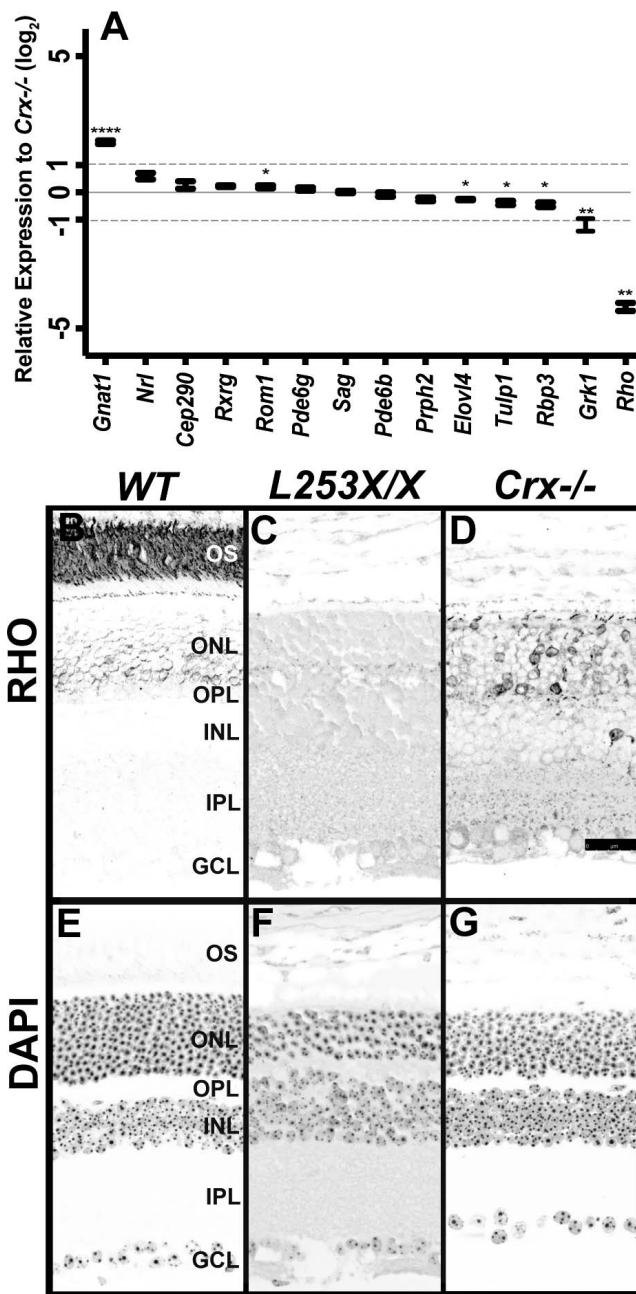
Other class III mutations show a greater impact on retina morphology and function than seen in *Crx*<sup>−/−</sup> mice.<sup>13,15,20</sup> The original report noted that degeneration was more rapid in *L253X/X* than *Crx*<sup>−/−</sup> retinas, but suggested differences in mouse background might be a factor.<sup>16</sup> To directly compare the impact of the *L253X* mutation with that of complete loss of *Crx*, we collected retinas from 1- and 3-month WT, *L253X/X*, and *Crx*<sup>−/−</sup> mice (all backcrossed 10 generations onto C57BL/6j) and assessed morphologic changes in H&E-stained sagittal sections through the optic nerve head (Figs. 2D–K). At 1 month, both *L253X/X* and *Crx*<sup>−/−</sup> retinas lacked photoreceptor outer segments (OS) and had thinner outer nuclear layers (ONL) than the WT control (Figs. 2E, 2F versus 2D). However, thinning was more pronounced in *L253X/X* than *Crx*<sup>−/−</sup>: *L253X/X* retina had only four to five rows of ONL nuclei, while *Crx*<sup>−/−</sup> maintained 7 to 8 rows at the same age (Fig. 2E versus 2F). Quantitative morphometry measures showed that the ONL of *L253X/X* was significantly thinner than that of *Crx*<sup>−/−</sup> at multiple positions (Fig. 2G, black versus solid gray). By 3 months, both models had similarly degenerated retinas with only 2–3 rows of ONL cells left (Fig. 2I versus 2J; 2K). Since both mutants showed normal ONL thickness at P7 through P10 (Supplementary Figs. S2A, S2B, and Tran et al.<sup>13</sup>), these results suggest that ONL degeneration occurs earlier or at a faster pace in *L253X/X* mice than in *Crx*<sup>−/−</sup>.

In addition to progressive ONL loss, *L253X/X* also underwent age-dependent thinning of the outer plexiform layer (OPL; Figs. 2E, 2I, Supplementary Figs. S2D, S2I, S2N, and Won et al.<sup>16</sup>). In contrast, nonphotoreceptor cell layers (e.g., inner nuclear layer, ganglion cell layer) were largely unchanged in *L253X/X* compared to WT controls at various ages (Supplementary Figs. S2C–Q). These results suggest that the *L253X* mutation mainly affects photoreceptor structural integrity and survival.

### *L253X/X* Mice Exhibit Photoreceptor Gene Misregulation Distinct From *Crx*<sup>−/−</sup>

To further examine if the more severe morphologic phenotype of *L253X/X* retinas reflected greater changes in CRX target gene expression, we collected RNA from *L253X/X* and *Crx*<sup>−/−</sup> retinas at P10 during photoreceptor terminal differentiation prior to cell death.<sup>13</sup> Transcripts of 14 known CRX-dependent genes were quantified by qRT-PCR and compared both with WT levels (Supplementary Fig. S2R) and between the two models (Fig. 3A). Although most of these genes were dysregulated in both mutants compared to WT mice (Supplementary Fig. S2R), half of them showed statistically significant expression differences between *L253X/X* and *Crx*<sup>−/−</sup> (Fig. 3A).<sup>13,21</sup> Three of these, *Gnat1*, *Grk1*, and especially *Rho* differed markedly (>2-fold) between the two models (Fig. 3A). Such differences in essential photoreceptor gene expression likely contribute to the morphologic and phenotype differences between the two models.

To confirm qRT-PCR results at the protein level, we performed immunofluorescent staining for RHO on retinal sections of 1-month-old *L253X/X*, *Crx*<sup>−/−</sup> and WT control mice. RHO was chosen because it displayed the largest difference in the RNA level between *L253X/X* and *Crx*<sup>−/−</sup>. As expected, in WT retina, RHO was predominantly localized to the rod outer segments (OS) with faint staining seen in ONL cell bodies (Fig.



**FIGURE 3.** *L253X/X* has measurable gene expression differences relative to *Crx*<sup>−/−</sup>. (A) Expression at P10 of the indicated photoreceptor genes was measured by qRT-PCR and compared between *L253X/X* and *Crx*<sup>−/−</sup> at P10. The gray dashed lines mark 2-fold differences (±1 log<sub>2</sub> relative expression) between the two genotypes (mean ± SEM,  $n \geq 3$ ; \*  $P < 0.05$ , \*\*  $P < 0.01$ , \*\*\*\*  $P < 0.0001$ ; unpaired *t*-test with Welch's Correction). (B–D) RHO immunostaining of retinal sections of the indicated mice at 1 mo revealed less RHO expression in *L253X/X* retinas than *Crx*<sup>−/−</sup> (signal in black, scale bar: 25 μm). (E–G) DAPI counterstaining of the above sections (signal in black).

3B). In contrast, *L253X/X* retina displayed no detectable RHO immunostaining (Fig. 3C), while *Crx*<sup>−/−</sup> retina still had weak and spotty RHO signals localized to individual ONL cell bodies and inner segments (Fig. 3D). These results supported the RNA expression data and suggest that *L253X* mutant protein negatively affects transcription.

Overall, the comparison between *L253X/X* and *Crx*<sup>−/−</sup> indicates that *L253X/X* disease is more severe than that



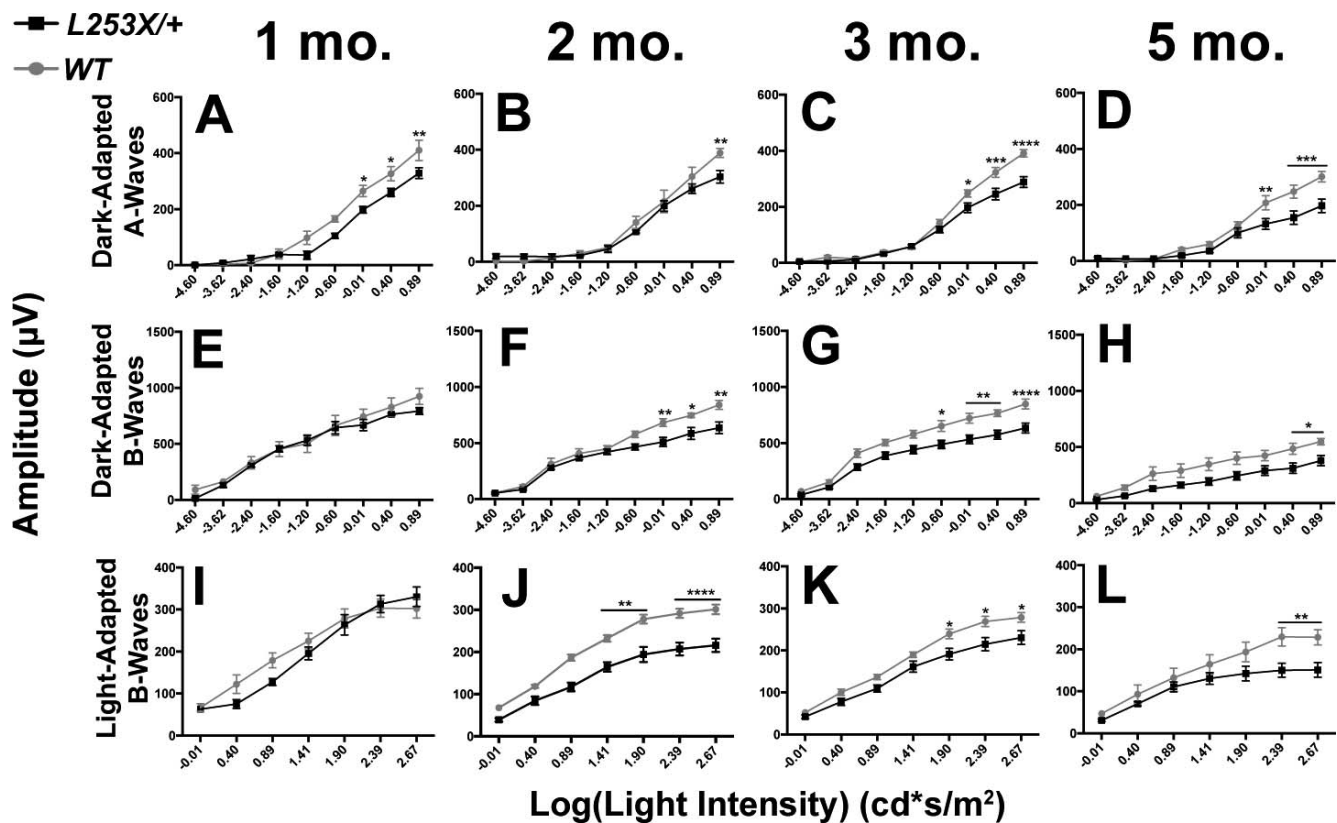


FIGURE 4. *L253X/+* retinas exhibit reduced rod and cone light responses. ERG responses of *L253X/+* mice (black lines) and WT (gray lines) were tested at four adult ages, and dark-adapted and light-adapted A- and B-wave amplitudes quantified: (A, E, I) 1 mo; (B, F, J) 2 mo; (C, G, K) 3 mo; and (D, H, L) 5 mo (mean  $\pm$  SEM,  $n \geq 3$ ; \*  $P < 0.05$ , \*\*  $P < 0.01$ , \*\*\*  $P < 0.0006$ , \*\*\*\*  $P < 0.0001$ ; 2-way ANOVA with Sidak's multiple comparisons test).

resulting from loss of *Crx*, supporting a dominant effect of the mutation.<sup>8,13</sup> The more severe photoreceptor deficits in *L253X/X* thus presumably result from antimorphic activity of the truncated CRX protein.

### Heterozygous *L253X/+* Mice Show Mild Decreases in Rod and Cone Function

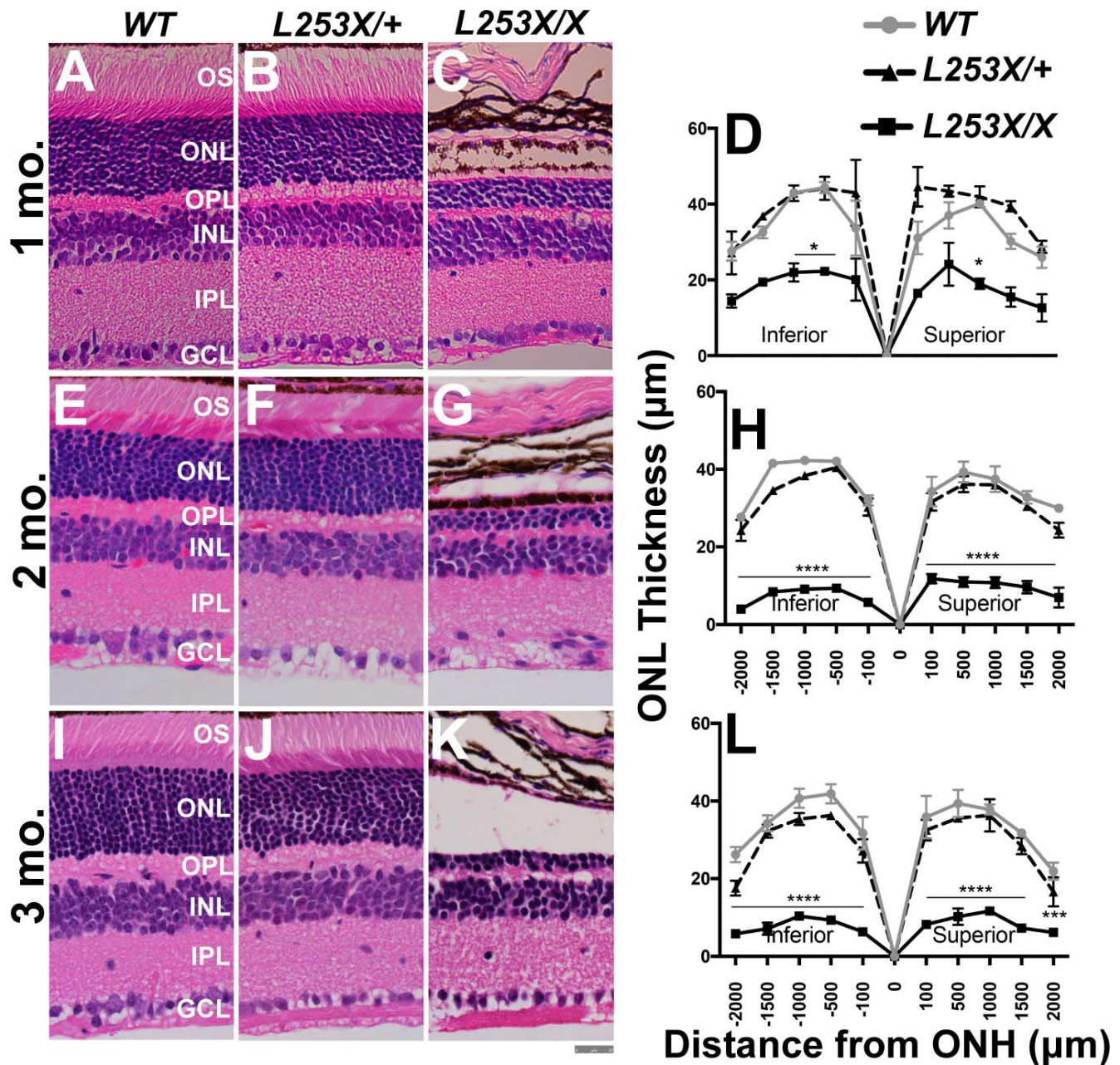
The *L253X* (*TVRM65*) mutation was originally reported to cause recessive retinal disease.<sup>16</sup> However, if the mutant protein possesses antimorphic activity, we would hypothesize that one copy would produce a photoreceptor phenotype. To test this, we investigated photoreceptor function in *L253X/+* retinas by assessing electrical responses to whole retina light stimulation. ERGs recorded at 1, 2, 3, and 5 months revealed *L253X/+* mice exhibit mild but significant reductions in ERG amplitudes compared to WT mice (Fig. 4). At all ages tested, dark-adapted A-waves were reduced in *L253X/+* mice, especially at high stimulus intensities (Figs. 4A–D, black line versus gray line). Dark-adapted B-waves of *L253X/+* mice were comparable to WT at 1 month (Fig. 4E), but significantly reduced at 2, 3, and 5 months (Figs. 4F–H, black line versus gray line). The degree of A-wave or B-wave reductions appeared to be constant, suggesting that the defects were unlikely a result of progressive degeneration. Reduction in cone function was also observed: Light-adapted B-waves were significantly decreased in *L253X/+* mice at 2, 3 and 5 months (Figs. 4J–L), but not at 1 month (Fig. 4I). These results suggest that *L253X/+* mice have deficits in both rod and cone function, resembling a mild retinopathy.

### *L253X/+* Retinas Do Not Degenerate

To test whether the functional defects correspond to morphologic changes, we measured retinal thickness in H&E stained retinal sections, comparing *L253X/+* and *L253X/X* with WT mice at three ages (Fig. 5). Despite the progressive retinal degeneration seen in *L253X/X* mice (Figs. 5C, 5D, 5G, 5H, 5K, 5L; Supplementary Figs. S2C–Q), *L253X/+* showed no degeneration or morphologic abnormalities compared to WT controls up to 3 months of age (Figs. 5B, 5D, 5F, 5H, 5J, 5L; Supplementary Figs. S2C–Q).

### *L253X/+* Retinas Display Dynamic Changes of Photoreceptor Gene Expression

To examine developmentally regulated photoreceptor gene expression in *L253X/+* versus WT mice, we performed qRT-PCR analyses at two early ages: P10, when rods/cones are still terminally differentiating, and 1 month when rods/cones are fully mature in normal retinas. The genes tested were previously shown to dramatically change in expression during WT rod and cone development.<sup>22</sup> Figure 6A shows expression changes of these genes in *L253X/+* relative to WT mice at both ages. Gene expression conformed to three general trends in addition to the slight increase in total *Crx* expression (dashed black line) noted above: *Gnat1*, *Rbo*, and *Sop* (solid black lines), markers of mature photoreceptors, were significantly decreased in *L253X/+* at P10 but attained WT levels by 1 month. *Rxrg* (dotted gray line), which in WT retina loses expression over development,<sup>22</sup> was elevated in *L253X/+* at P10 but decreased closer to WT



**FIGURE 5.** *L253X/+* retinas appear morphologically similar to *WT* up to 3 mo. H&E-stained retinal sections from *L253X/+*, *L253X/X*, and *WT* mice at the indicated ages: 1 mo (**A–C**); 2 mo (**E–G**); and 3 mo (**I–K**) were imaged at  $\times 40$  magnification (scale bar: 25  $\mu$ m). Images were taken approximately 500  $\mu$ m from the ONH, where differences in layer thickness are less significant. (**D**, **H**, **L**) ONL thickness was quantified by morphometry at five set distances (in microns) on either side of the ONH (mean  $\pm$  SEM,  $n \geq 3$ ; \*  $P < 0.05$ , \*\*  $P < 0.01$ , \*\*\*  $P < 0.001$ , \*\*\*\*  $P < 0.0001$ ; 2-way ANOVA with Tukey's multiple comparisons test).

levels by 1 month. The other genes tested, which increase over development in *WT* retina, showed similar relative expression levels at both ages that did not vary considerably from *WT* (solid gray lines). Together, these gene expression trends suggest that *L253X/+* retinas exhibit altered gene expression early in photoreceptor development, similar to other *Crx* mutant models.<sup>21</sup>

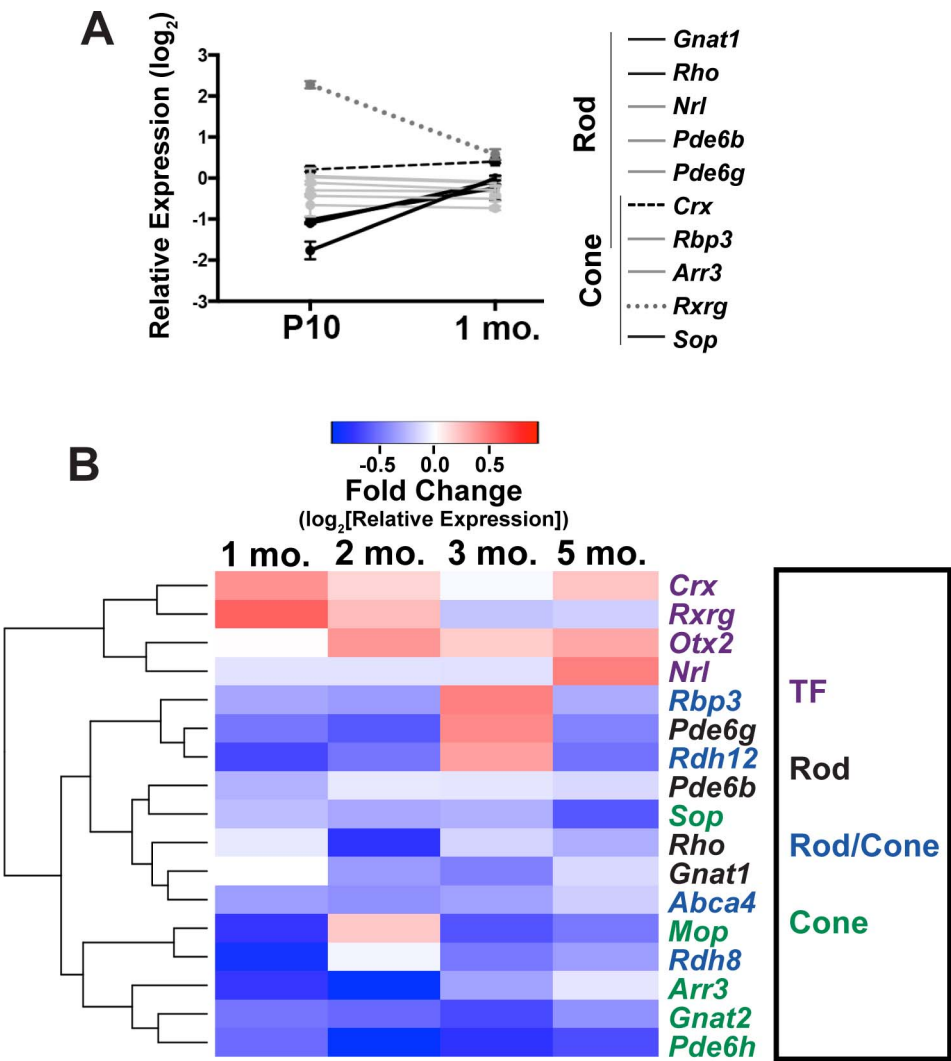
To determine if *L253X/+* retinas maintain photoreceptor gene expression after differentiation, we used qRT-PCR to examine expression of 17 essential rod and cone genes in *WT* and *L253X* mutants at other adult ages. The results at 1, 2, 3 and 5 months (presented in a heatmap in Fig. 6B) suggest a complex phenotype. Genes coding for transcription factors

*Crx*, *Rxrg*, *Otx2*, and *Nrl* were upregulated at many of the time points tested. The expression of essential rod and cone phototransduction genes showed a less clear pattern across ages, but in general all exhibit phases of reduced expression relative to *WT*.

#### Late PTC-Caused C-Terminal Truncation Reduces Transactivation Function of CRX

As an initial step to understand the molecular mechanism underlying misregulation of gene expression in *L253X* mutant retinas, we asked whether a short C-terminal truncation caused by a late PTC affects the ability of CRX to transactivate





**FIGURE 6.** *L253X/+* retinas display dynamic changes in photoreceptor gene expression. qRT-PCR analyses of *L253X/+* expression levels (relative to *WT* mice) of the indicated rod/cone genes are shown at P10 and 1 mo (mean  $\pm$  SEM,  $n \geq 3$ ). Lines connecting expression levels of each gene at the two ages show the dynamic changes in expression, and reveal four different expression trends indicated here by line color and solidity, as described in Results. **(B)** Heatmap depicting qRT-PCR-detected expression changes of the listed photoreceptor genes in *L253X/+* relative to *WT* at the indicated ages. Data are clustered based on gene expression patterns (log<sub>2</sub> [mean fold-change],  $n \geq 3$ ). Colored gene names indicate expression in specific photoreceptor subtype(s). TF indicates transcription factors expressed by rods and/or cones that are essential for cell fate specification.

target gene expression. This was tested using transient transfection assays in HEK293T cells with a Rhodopsin (*Rho*) promoter-luciferase reporter (*BR130-Luc*).<sup>12</sup> Vectors expressing either the full-length (normal) or truncated CRX<sup>L253X</sup> were cotransfected with an NRL vector to measure their ability to transactivate the *BR130-Luc* reporter. CRX<sup>L253X</sup> did show some dose-dependent ability to activate the *Rho* promoter, suggesting that it does bind DNA and interacts with NRL, but the maximum activity was greatly reduced compared to the normal CRX protein (~30% of normal activity; Fig. 7).

## DISCUSSION

### *L253X* Mouse Provides a New Class III Model for Mild Dominant Cone-Rod Dystrophy

Although *L253X* (*TVM65*) was originally identified as a recessive *Crx* mutation,<sup>16</sup> several pieces of evidence suggest that *L253X* is antimorphic with dominant inheritance. First,

*L253X* retinas overexpress the mutant gene product, a hallmark of class III mutations that amplifies the mutation's effects. Second, the photoreceptors in homozygous *L253X/X* mice degenerate earlier than *Crx*<sup>-/-</sup>, and show patterns of photoreceptor gene dysregulation that are distinct from *Crx*<sup>-/-</sup>. Thus, the presence of the mutant *L253X* protein causes more severe pathology than absence of CRX altogether, which is consistent with the antimorphic effect of other class III mutations<sup>11,13,15</sup>. Furthermore, heterozygous *L253X/+* mice have early gene expression differences that largely resolve by 1 month, suggesting that full maturation may be delayed in these retinas. Although gene expression changes in adults were modest, rod and cone functional deficits were detectable as early as 1 month. Previous work on class III mutants has emphasized the effects that small changes in gene expression over a large number of genes can have on photoreceptor function.<sup>21</sup> Collectively, these findings identify *L253X* as a class III *Crx* mutation with the phenotypes summarized in the Table.

Compared to previously reported class III mutations, pathogenicity of *L253X* is relatively mild. Unlike *E168d2/+*

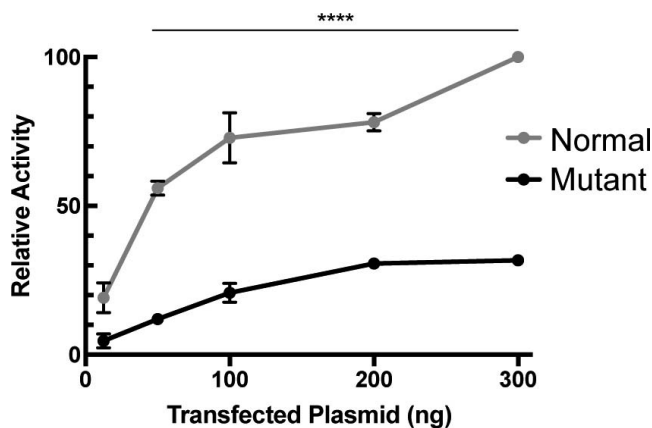


FIGURE 7. L253X retains minimal ability to transactivate rhodopsin promoter in the presence of NRL. Dual luciferase assays showing a dose-dependent increase in CRX transactivation activity of normal or mutant ( $CRX^{L253X}$ ) in the presence of constant amount of NRL (mean  $\pm$  SEM,  $n = 3$ ; \*\*\*\*  $P < 0.0001$ ; 2-way ANOVA with Bonferroni's multiple comparisons test).

photoreceptors, which lose their OS and undergo progressive degeneration,<sup>13</sup> L253X/+ photoreceptors show normal morphology and do not degenerate. The ERG deficits do not worsen with age relative to C57BL/6J controls. The gene expression changes are also relatively modest with a multiphasic nature. This evolving expression pattern with increased levels of many TFs, but modest depletion of phototransduction cascade components, likely reflects an inability of the cells to maintain a proper homeostasis. This data would suggest that unknown feedback mechanisms are operational but unable to achieve the necessary precision. In adults, L253X/+ photoreceptors are alive and functional through 5 months, but it remains to be determined if the phenotype worsens at older ages or if the retina is more susceptible to environmental insults as reported with other class III mutants. Nevertheless, given the huge phenotypic variability of human CRX diseases, the L253X mouse provides a valuable animal model for understanding the pathogenesis of class III mutations and the underlying molecular mechanisms that determine phenotype severity, age of onset and disease progression.

### The Molecular Mechanism(s) Underlying L253X Pathogenicity

The pathogenicity of class III mutations is generally determined by two factors, the activity and dose of the mutant protein. L253X protein contains an intact DNA binding domain but lacks a portion of the C-terminal activation domain (Fig. 1A). Thus, it likely retains the ability to bind DNA, but loses some aspect of transactivation function. Transient transfection assays showed that  $CRX^{L253X}$ , a truncated CRX protein similar to L253X, retained only 30% of the normal CRX activity, very similar to the published reports.<sup>12,14</sup> By comparison, the class III mutant protein E168d2 showed 25% to 30% of the

transactivation activity under similar conditions.<sup>12,13</sup> Previous reports have also shown  $CRX^{L253X}$  binds appropriate DNA sequence elements very similar to full-length CRX.<sup>14</sup> It is conceivable that, as previously reported for E168d2,<sup>13</sup> L253X protein may compete with normal CRX to bind to target genes, interfering with normal CRX function in heterozygous mice. However, transient transfection assays testing this hypothesis did not show this effect on the *Rbo*-promoter luciferase reporter (data not shown). The antimorphic competition may not be limited to CRX, but could also affect other homeodomain transcription factors that bind to similar DNA targets, and as such transient cell transfection assays on a single reporter that lacks the native chromatin context and regulatory architecture may not be the proper experiment. Supporting this possibility, the retinas of L253X/X mutants, like E168d2/d2,<sup>13</sup> showed more severe degeneration and biochemical defects than  $Crx^{-/-}$  retinas. Other potential targets could include OTX2,<sup>25</sup> which is upregulated in L253X retinas (Fig. 6B). Additional experiments are required to demonstrate dominant-negative activity of L253X on CRX and OTX2. Despite the similarity between L253X and E168d2, L253X/+ mice have a remarkably milder phenotype than E168d2/+; it is unlikely that the <10% difference in transactivation activity can account entirely for the significant difference between the two models. We therefore propose that the relative amounts of mutant and normal CRX protein must also play a role in phenotype severity.

### L253X Allele-Specific Overproduction of Mutant mRNA and Protein Has Implications for the PTC Position Effect

Careful analysis of *Crx* mRNA and protein in L253X/+ and L253X/X retinas clearly shows an increased ratio of mutant/normal protein and mRNA. This finding supports the hypothesis that, like other class III models, L253X retinas accumulate mutant transcripts, leading to overproduction of the CRX mutant protein. Interestingly, however, the degree of mutant RNA accumulation in L253X/+ is lower than that of the other class III models carrying early PTC mutations. L253X mice only display a modest accumulation (57% mutant/43% normal), especially when compared to that observed in E168d2/+ mice at the same age (85% mutant/15% normal).<sup>13</sup> Also, when considered with the quantification of total *Crx* mRNA, the L253X/+ must result in a downregulation of the normal allele as there is no aggregate overproduction of *Crx*. This suggests that the L253X mRNA is still "sensed" by the feedback regulatory mechanism, which functions normally in the mutant retina. Future studies could probe this feedback mechanism by comparing to the E168d2/+ retinas where although there is a gross increase in total *Crx* mRNA, the normal allele maintains normal expression levels.<sup>13</sup> Both mouse models show overproduction of mutant protein relative to WT, but L253X accumulates at a slower rate than E168d2.

The mechanism for allele-specific overproduction of mutant *Crx* in class III models is not well understood, but likely stems from increased RNA stability due to the presence of the PTC in

TABLE. Summary of L253X Related Phenotypes

Phenotype	<i>Crx-L253X/X</i>	<i>Crx-L253X/+</i>
Morphology (histology)	ONL degeneration earlier than $Crx^{-/-}$	No detectable abnormality up to 5 mo
Function (ERG)	Lack of rod and cone responses to light	Minor reductions in rod and cone function
Gene expression (qRT-PCR)	Altered gene expression distinct from $Crx^{-/-}$	Dynamic changes from WT
CRX protein/RNA levels	L253X protein/RNA overproduction	Increased ratio of L253X versus normal

the last exon. This hypothesis further predicts that the position of the PTC influences the rate of mutant RNA accumulation. Indeed, *L253X* with a later PTC showed less RNA accumulation than the two early PTC alleles, *E168d2* and *A182d1* (Fig. 1). Thus, the earlier the PTC is, the more mutant product accumulates. These differences between early versus late PTCs may be attributed to the length of PTC-determined 3'UTR, within which multiple discrete elements could independently and additively contribute to mRNA hyperstability. Future identification of novel RNA regulatory elements shared among different PTC mutations may shed light on the mechanisms for mutant transcript hyperstability and phenotype variability in class III *CRX* disease. It is also notable that the accumulation of mutant protein in *L253X* retinas is more prominent than mutant RNA, raising the new possibility that altered protein turnover rate could also contribute to mutant protein accumulation.

In conclusion, our in-depth characterization of the *Crx-L253X* (*TVRM65*) mouse supports the classification of *L253X* as a dominant class III *Crx* mutation and provides a new animal model for understanding *CRX*-associated dominant retinopathies. The insights gained will hopefully lead to new treatment strategies for this complex disease.

### Acknowledgments

The authors thank Connie Cepko for providing *Crx*<sup>-/-</sup> mice, Guangyi Ling, and Belinda Dana of the Immunology core for technical assistance in histology and immunohistochemistry assays, and Mingyan Yang for mouse genotyping and other technical assistance. The authors thank Brandon McKethan of Bio-Rad Laboratories for designing ddPCR assays to quantify mutant versus normal transcripts in *L253X*/+ retinas.

Supported by National Institutes of Health grants EY012543 and EY025272 (SC); EY013360 (Washington University [WU]) and EY002687 (WU, Department of Visual Science [WU-DOVS]); and unrestricted funds from Research to Prevent Blindness (WU-DOVS).

Disclosure: **P.A. Ruzycski**, None; **C.D. Linne**, None; **A.K. Hennig**, None; **S. Chen**, None

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