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Thyroid hormone receptors mediate two distinct mechanisms of long-wavelength vision

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Thyroid hormone (TH) signaling plays an important role in the regulation of long-wavelength vision in vertebrates. In the retina, thyroid hormone receptor β (*thrb*) is required for expression of long-wavelength-sensitive opsin (*lws*) in red cone photoreceptors, while in retinal pigment epithelium (RPE), TH regulates expression of a cytochrome P450 enzyme, *cyp27c1*, that converts vitamin A₁ into vitamin A₂ to produce a red-shifted chromophore. To better understand how TH controls these processes, we analyzed the phenotype of zebrafish with mutations in the three known TH nuclear receptor transcription factors (*thraa*, *thrab*, and *thrb*). We found that no single TH nuclear receptor is required for TH-mediated induction of *cyp27c1* but that deletion of all three (*thraa*^{-/-}; *thrab*^{-/-}; *thrb*^{-/-}) completely abrogates its induction and the resulting conversion of A₁- to A₂-based retinoids. In the retina, loss of *thrb* resulted in an absence of red cones at both larval and adult stages without disruption of the underlying cone mosaic. RNA-sequencing analysis revealed significant downregulation of only five genes in adult *thrb*^{-/-} retina, of which three (*lws1*, *lws2*, and *miR-726*) occur in a single syntenic cluster. In the *thrb*^{-/-} retina, retinal progenitors destined to become red cones were transfated into ultraviolet (UV) cones and horizontal cells. Taken together, our findings demonstrate cooperative regulation of *cyp27c1* by TH receptors and a requirement for *thrb* in red cone fate determination. Thus, TH signaling coordinately regulates both spectral sensitivity and sensory plasticity.

thyroid hormone | retinal development | cone photoreceptors | vitamin A₂ | zebrafish

The spectral sensitivity of a visual pigment is determined by the amino acid sequence of the opsin protein and the chemical structure of its covalently bound, vitamin A-based chromophore (1, 2). In the course of evolution, opsin gene duplication followed by the divergence of amino acid sequences involved in spectral tuning has resulted in a multiplicity of opsins with peak spectral sensitivities ranging from UV to red (3). For example, zebrafish possess four cone photoreceptor subtypes that each express a different opsin: UV cone opsin (*sws1*), blue cone opsin (*sws2*), green cone opsin (*rh2*), and red cone opsin (*lws*) (Fig. 1A). The zebrafish genome contains two red cone opsin paralogs (*lws1* and *lws2*) and four green cone opsin paralogs (*rh2-1*, *rh2-2*, *rh2-3*, and *rh2-4*) (4, 5).

The spectral sensitivities of photoreceptors appear to be stable across the lifespan of most avian and mammalian species (3). In contrast, numerous species of aquatic vertebrates are capable of dynamically tuning their spectral sensitivities at different stages of the life cycle or in response to changing environmental conditions (6, 7). For example, when salmon or lamprey migrate from the blue-green waters of the open ocean to the red-shifted spectral milieu of inland streams to spawn, they shift the spectral sensitivity of their visual system to match that of the new environment (8–10). This shift is primarily mediated by the expression of a cytochrome P450 enzyme, *cyp27c1*, which converts

vitamin A₁ into vitamin A₂ to produce a red-shifted chromophore in the retinal pigment epithelium (RPE) (Fig. 1A) (9, 11). Replacement of the A₁-based chromophore, 11-*cis* retinaldehyde, with the A₂-based chromophore, 11-*cis* 3,4-didehydroretinaldehyde red-shifts the sensitivity of red cone opsins by nearly 60 nm (11, 12). Conversely, the A₁-to-A₂ shift has only a minimal effect on the spectral sensitivity of short wavelength-tuned opsins (i.e., UV- and blue-sensitive opsins) (11, 12). Therefore, the primary effect of *cyp27c1* expression is to extend the sensitivity of the longest wavelength-sensitive opsin further into the near infrared, thereby facilitating vision in turbid waters where longer wavelength light predominates. Despite the importance of this mechanism of sensory plasticity in aquatic vertebrates, the transcription factors that control *cyp27c1* expression are currently unknown. One clue to the mechanism of *cyp27c1* regulation is that exogenous application of TH induces an A₁-to-A₂ shift in the eyes of various teleost species (7, 12). This finding suggests a role for TH signaling in regulating *cyp27c1* expression.

TH signaling has also been shown to regulate opsin expression in the retina. The application of TH drives both the transition from UV cone opsin (*sws1*) to blue cone opsin (*sws2*) expression and the induction of a longer-wavelength-sensitive (red-shifted) green cone opsin (*rh2*) paralog in salmonid species (13, 14).

Significance

Spectral sensitivity is determined by the components of the light-sensitive visual pigment: an opsin protein and a covalently bound chromophore. Thyroid hormone signaling has been shown to regulate both the expression of long-wavelength (red) opsin and *cyp27c1*, an enzyme that converts the vitamin A₁-based chromophore to the red-shifted vitamin A₂-based form. Here, we show that all three zebrafish thyroid hormone nuclear receptors play a role in mediating induction of *cyp27c1* expression in response to thyroid hormone and that mutations in thyroid hormone receptor β result in an absence of red cones and the transfating of red cone precursors into UV cones and horizontal cells in zebrafish. These results demonstrate that thyroid hormone receptors regulate two distinct aspects of long-wavelength vision.

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The authors declare no competing interest.

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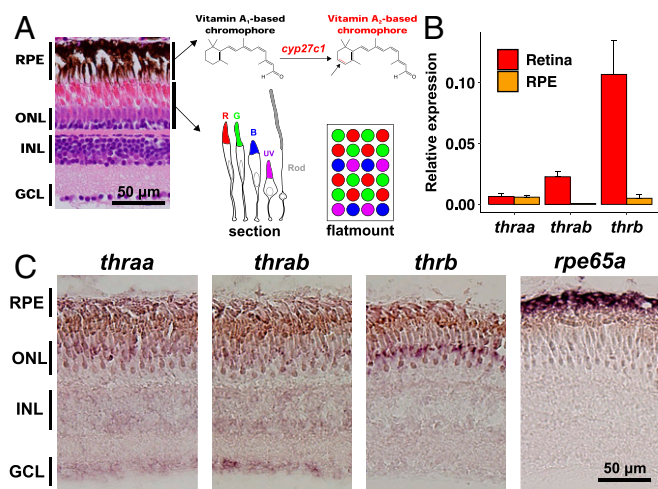


Fig. 1. TH receptor expression in zebrafish retina and RPE. (A) Hematoxylin and eosin-stained section of 9-mo-old zebrafish eye. The accompanying diagram shows *cyp27c1*-mediated conversion of 11-*cis* retinaldehyde (the vitamin A₁-based chromophore) into 11-*cis* 3,4-didehydroretinaldehyde (the vitamin A₂-based chromophore) in the RPE, and photoreceptor subtypes in the ONL. (B) Quantification of TH receptor expression in the retina and RPE of 5-mo-old WT zebrafish by qPCR. Relative expression (normalized to *rp13a* expression [$2^{-\Delta\Delta Ct}$]; mean \pm SD; $n = 3$ for all transcripts, except $n = 1$ for *thrab* in RPE due to lack of detectable transcript in two samples). (C) ISH of TH receptors in 9-mo-old *albino* zebrafish eye demonstrates that all TH receptors are expressed in the retina, whereas there is no detectable signal in the RPE. ISH with a probe against *rpe65a*, an RPE-specific gene, confirms the presence of RPE in the histologic sections.

Recent studies have shown that TH drives the expression of red-shifted green and red cone opsin paralogs in zebrafish as well (15). There is also evidence that *thrb* regulates the expression of red cone opsins and represses the expression of UV cone opsin orthologs in zebrafish, mouse, and human retinal organoids (16–18). Thus, TH signaling is likely to play an evolutionarily conserved role in the regulation of red cone opsin expression in vertebrates, thereby facilitating long-wavelength vision.

To elucidate the mechanisms by which TH receptors regulate long-wavelength vision, we analyzed the phenotype of zebrafish with mutations in the three known TH nuclear receptors: *thraa*, *thrab*, and *thrb*. We found that all three zebrafish TH receptors play a role in driving *cyp27c1* expression in RPE and that mutations in *thrb* result in an absence of red cones and transfating of red cone precursors into UV cones and horizontal cells in larvae. These results show how TH signaling mediates both spectral sensitivity and sensory plasticity.

Results

***thraa*, *thrab*, and *thrb* Redundantly Regulate *cyp27c1* Expression and the Vitamin A₁-to-A₂ Switch.** The zebrafish genome encodes three TH receptors: *thraa*, *thrab*, and *thrb* (19). To begin to evaluate the role of these receptors in zebrafish retina and RPE, we measured their expression by quantitative RT-PCR (qPCR). We found that all three receptors are expressed in the adult retina (Fig. 1B and *SI Appendix*, Fig. S1A). In RPE, in contrast, *thraa* and *thrb* are expressed while *thrab* is barely detectable (Fig. 1B and *SI Appendix*, Fig. S1A).

Next, we determined the cellular expression pattern of TH receptors by performing in situ hybridization (ISH) on adult *albino* zebrafish eyes (20). We used the *albino* strain to permit visualization of gene expression in RPE, which is heavily pigmented with melanin in wild-type (WT) fish. In the retina, we found *thraa* and *thrab* to be weakly expressed in all three layers (ONL, outer nuclear layer; INL, inner nuclear layer; and GCL,

ganglion cell layer), whereas *thrb* expression was restricted to the ONL (Fig. 1C). The expression of all three receptors in the ONL appears to be restricted to a subset (or subsets) of photoreceptors, but the weakness of the *thraa* and *thrab* signals makes definitive assessment difficult. Publicly available single cell RNA-sequencing (RNA-seq) data from adult zebrafish retina confirm that *thraa* and *thrab* are broadly expressed across multiple retinal cell types, whereas *thrb* is specifically expressed in cones (21). In RPE, we were unable to detect any signal above background for any of the TH receptors by ISH (Fig. 1C). A strong signal for *rpe65a*, an RPE-specific transcript, confirmed the presence of RPE in our histologic sections (Fig. 1C).

To determine which TH receptors mediate TH-driven induction of *cyp27c1* expression and the resultant vitamin A₁-to-A₂ switch, we measured these processes in TH receptor mutant zebrafish. For this purpose, we utilized previously published zebrafish lines with mutations in *thraa* and *thrab* (22). In addition, we used CRISPR-Cas9 technology to engineer two *thrb* mutant alleles: *thrb*^{sl627} (which contains a 25-bp deletion in exon 9) and *thrb*^{sl628} (which contains a 52-bp deletion in exon 9). Both *thrb* mutant lines are predicted to result in a frame shift followed by a premature stop codon. These lines will both herein be referred to as *thrb*^{-/-}. All TH receptor mutations used in this study are predicted to truncate the receptor upstream of the ligand-binding domain (*Materials and Methods* and *SI Appendix*, Fig. S2). To evaluate the role of individual TH receptors in the TH-mediated induction of *cyp27c1* expression, we maintained *thraa*^{-/-}, *thrab*^{-/-}, and *thrb*^{-/-} zebrafish and WT control fish in water containing TH (300 μ g/L L-thyroxine) or vehicle for 3 wk. We then used qPCR to quantify *cyp27c1* expression and high-performance liquid chromatography (HPLC) to measure the relative amounts of A₁ and A₂ retinaldehydes. TH treatment resulted in robust induction of *cyp27c1* expression and a marked increase in A₂ retinaldehyde content in all three mutants and WT controls (Fig. 2A–F). Although no reduction in *cyp27c1* expression was observed in TH-treated mutants, there was a statistically significant decrease in A₂ content in TH-treated *thraa*^{-/-} fish relative to vehicle-treated mutants (Fig. 2A and D). *thrab*^{-/-} and *thrb*^{-/-} fish displayed normal *cyp27c1* induction and A₂ content (Fig. 2B, C, E, and F). Thus, no single TH receptor is required for induction of *cyp27c1* expression and A₁-to-A₂ conversion.

To assess whether the various TH receptors might act redundantly, we intercrossed the three mutant lines to create all possible double and triple mutant combinations. We then exposed the fish to either TH or vehicle and measured *cyp27c1* expression and A₁ and A₂ content. All double mutant fish showed reduced A₂ content (Fig. 2K–M), whereas only *thraa*^{-/-};*thrb*^{-/-} fish displayed a statistically significant decrease in *cyp27c1* induction (Fig. 2G–I). Despite the relatively severe reduction in A₂ content in *thraa*^{-/-};*thrb*^{-/-} fish, there appeared to be some residual induction of *cyp27c1* expression in this mutant relative to vehicle controls (Fig. 2I). Only in *thraa*^{-/-};*thrab*^{-/-};*thrb*^{-/-} fish that lack all functional TH receptors, was there a complete failure of *cyp27c1* induction and A₂ production in response to TH treatment (Fig. 2J and N). Thus, all three TH receptors appear to contribute to TH-mediated induction of *cyp27c1* expression and A₂ retinoid production, with *thraa* and *thrb* appearing to be most important.

Red Cones Are Absent from *thrb*^{-/-} Retinas. A prior study demonstrated loss of red cone opsin expression in larval zebrafish using morpholinos targeting *thrb* (16). However, morpholino-induced phenotypes sometimes differ from those of the corresponding mutants, and the action of morpholinos is limited to the early larval stage (23). It also remains unclear whether the *thrb* morpholino phenotype is due to a simple loss of opsin expression or a defect in red cone fate determination. To address these questions, we analyzed the retinal phenotype of *thrb*^{-/-} fish at larval

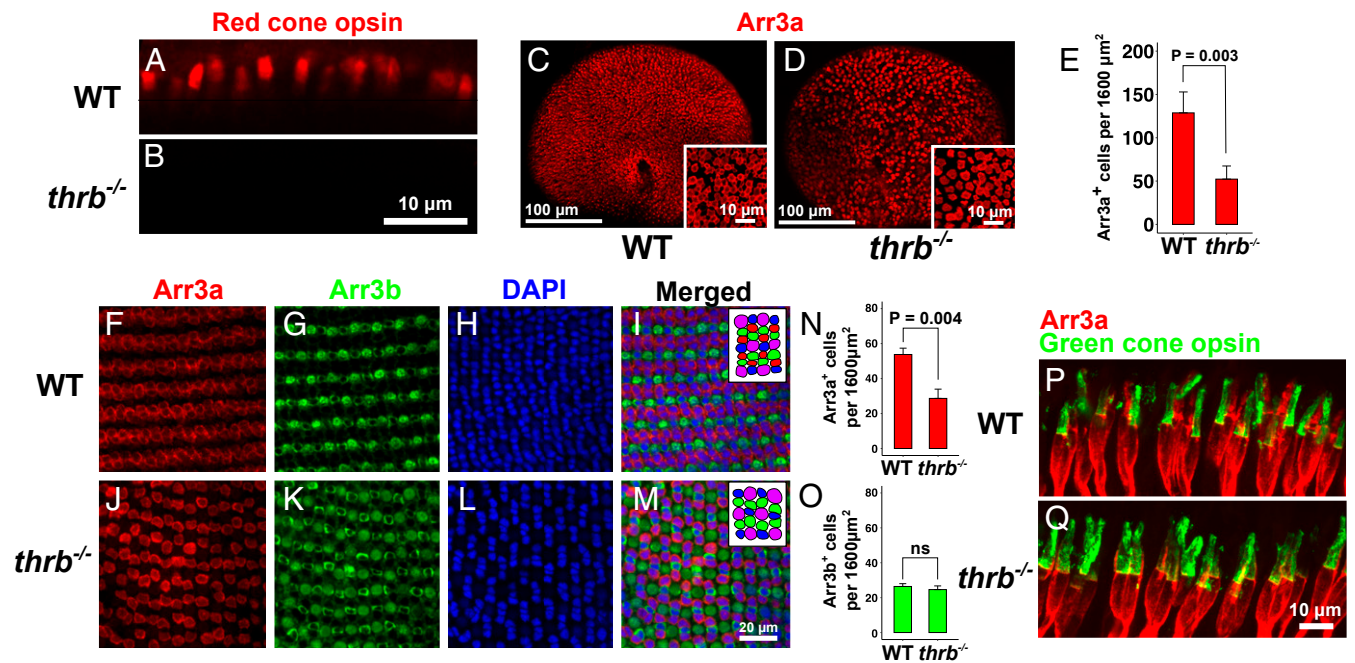


Fig. 3. Red cones are absent from larval and adult *thrb*^{-/-} retina. (A and B) Images of outer segments of 3 dpf larvae demonstrate a lack of red cone opsin⁺ cells in the 3 dpf *thrb*^{-/-} retina. (C and D) Wholemount views of 5 dpf WT and *thrb*^{-/-} retina stained with Arr3a antibody demonstrate a reduction in Arr3a⁺ cells in *thrb*^{-/-} retina. Insets are high-resolution images taken immediately dorsal to the optic nerve of separate clutch-matched larvae. (E) The number of Arr3a cells in the 5 dpf *thrb*^{-/-} retina is reduced relative to WT (mean ± SD; n = 4 per group). (F–M) Flatmount images of adult 5-mo-old WT and *thrb*^{-/-} retina stained with anti-Arr3a antibody (red and green cones), anti-Arr3b antibody (blue and UV cones), and DAPI (nuclei). (F and J) The double row of Arr3a⁺ cells in the WT retina is replaced with a single row in the *thrb*^{-/-} retina. (I and M) Combined images indicate that the patterning of the cone mosaic remains largely intact in the *thrb*^{-/-} retina. (N) Quantification of the number of Arr3a⁺ cells in the adult WT and *thrb*^{-/-} retina (mean ± SD; n = 3 per group) reveals that the total number of Arr3a⁺ cells is reduced by about half in the *thrb*^{-/-} retina. (O) Quantification of the number of Arr3b⁺ cells in the adult WT and *thrb*^{-/-} retina (mean ± SD; n = 3 per group) reveals that the number of Arr3b⁺ cells is unchanged. The reduction in the number of Arr3a⁺ cells in the *thrb*^{-/-} retina suggests that either red or green cones are absent. (P and Q) Sections of adult WT and *thrb*^{-/-} retina (inner and outer segments shown) stained with anti-Arr3a and anti-green cone opsin antibodies. In the WT retina, Arr3a⁺;green cone opsin⁺ cells (green cones) and Arr3a⁺;green cone opsin⁻ cells (red cones) are present. In the *thrb*^{-/-} retina, only Arr3a⁺;green cone opsin⁺ cells (green cones) are present, suggesting red cones are absent. All statistical analyses were performed using two-tailed t tests. *thrb*^{-/-} = *thrb*^{st1627/st1627} in A–O. *thrb*^{-/-} = *thrb*^{st1628/st1628} in P and Q. ns, not significant.

We next used RNA-seq to compare the transcriptome of the adult *thrb*^{-/-} retina to that of WT fish. In addition to *thrb* itself, we identify only five down-regulated genes (Fig. 5A and Dataset S1). Three of these genes reside in a single syntenic cluster and encode the two paralogous red cone opsins (*lws1* and *lws2*) and a microRNA (*miR-726*) (Fig. 5B). A homolog of this microRNA, which is of unknown function, was previously reported to be red cone-specific in medaka (*Oryzias latipes*) (27). Down-regulation of *thrb*, *lws1*, and *lws2* in *thrb*^{-/-} retina was additionally confirmed by qPCR (SI Appendix, Fig. S3). Examination of the distribution of RNA-seq reads at the *lws1/lws2/miR-726* locus indicates that the transcription of the immediately adjacent blue cone opsin gene (*sws2*) is unaffected by the *thrb* mutation (Fig. 5B).

Two other genes were found to be down-regulated in *thrb*^{-/-} retina: *si:busm1-57f23.1* and *mhc1uba* (Fig. 5A). The former is selectively expressed in developing eye and notochord and encodes a protein with similarity to the mammalian cysteine protease inhibitor cystatin (28). The expression pattern of *mhc1uba*, which encodes a component of the major histocompatibility complex, is less specific but includes the eye and other parts of the developing head (29). Only a single gene, of unknown function (*serinc2*), was found to be modestly up-regulated in *thrb*^{-/-} retina (Fig. 5A). Taken together, these findings indicate the presence of only very few red cone-specific genes in adult zebrafish. However, RNA-seq performed on whole retina may limit our ability to detect lowly expressed red cone-specific transcripts or those that are not exclusively red cone-specific.

Next, we conducted a preliminary evaluation of the distribution of cone photoreceptors in adult *thraa*^{-/-} and *thrab*^{-/-} retinas. Staining with anti-Arr3a, anti-Arr3b, and the nuclear marker DRAQ5, revealed a typical photoreceptor nuclear arrangement with a double row of Arr3a⁺ cells and a single row of Arr3b⁺ cells in both mutants (SI Appendix, Fig. S4). These results suggest the presence of a normal complement of the four cone photoreceptor subtypes. Given the broad expression pattern of *thraa* and *thrab* in multiple retinal layers (Fig. 1C), more subtle effects of these mutations on zebrafish retinal development may exist and will be the subject of future analyses.

Red Cone Precursors Are Transfated into UV Cones and Horizontal Cells in the *thrb*^{-/-} Retina. Morpholino-based knockdown of *thrb* in zebrafish larvae results in both a loss of red cone opsin⁺ cells and a corresponding increase in UV cone opsin⁺ cells (16). One possible explanation for this result is that red cone precursors are transfated into UV cones in the absence of *thrb*. To evaluate this possibility, we examined zebrafish carrying the *trp2:tdTomato* transgene, which consists of the tdTomato gene under control of the zebrafish *thrb* promoter (16). In these fish, tdTomato is first detected at 2 dpf in precursors that give rise to red cones, horizontal cells, and retinal ganglion cells (16). In WT retina at 3 dpf, we detected tdTomato exclusively in red cones (red cone opsin⁺ cells) in the ONL (Fig. 6A). In contrast, in *thrb*^{-/-} retinas, tdTomato⁺ cells occur both in the ONL and in the outermost part of the INL in a distribution similar to that of developing horizontal cells (30) (Fig. 6B). The tdTomato⁺ cells in the ONL

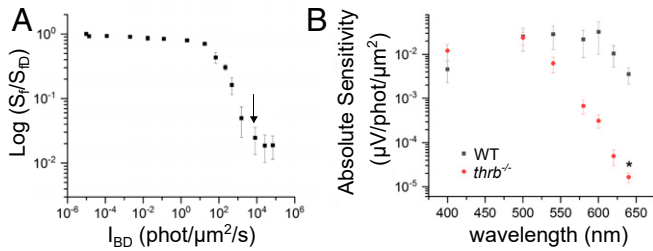


Fig. 4. Adult *thrb*^{-/-} fish exhibit diminished spectral sensitivity at long wavelengths. (A) Background light adaptation experiment used to determine the optimal background light intensity for saturating the rods and isolating the cone component of the retina response. Fractional flash sensitivity (S_i) measured by transretinal ERG recordings from WT zebrafish in 460-nm background light of increasing intensity, plotted normalized to the corresponding fractional flash sensitivity in darkness (S_{iD}). The initial gradual decline in dimmer backgrounds is driven by the light adaptation of the rods. Rod sensitivity declines steadily until a plateau is reached. Beyond this point, rod sensitivity becomes lower than cone sensitivity and the overall response from the retina is driven by the cones. We used the background intensity near the beginning of the plateau (8,109 phot/ $\mu\text{m}^2/\text{s}$; arrow) to suppress the rod responses and obtain transretinal ERG recordings of cone spectral sensitivity. (B) Absolute spectral sensitivity of cones obtained by transretinal ERG recordings in the background light identified in A. *thrb*^{-/-} fish (red circles; $n = 9$) lacked the long-wavelength spectral sensitivity (mean \pm SEM) observed in WT fish (black squares; $n = 8$). The difference in the absolute spectral sensitivity between the two groups was found to be statistically significant at 640 nm, indicating compromised red cone function in the *thrb*^{-/-} fish. All statistical analyses were performed using two-tailed *t* tests. * $P < 0.05$. *thrb*^{-/-} = *thrb*^{st1627/st1627}. Phot, photons.

of *thrb*^{-/-} retinas lack both red cone opsin expression and the presynaptic axonal narrowing seen in tdTomato⁺ cells in WT retinas (Fig. 6A and B). Furthermore, tdTomato⁺ cells in *thrb*^{-/-} retinas do not express Arr3a, a marker of mature red and green cones (Fig. 6D). We note that in the 3 dpf WT retina, Arr3a labeling is primarily in tdTomato⁺ red cones (Fig. 6C), which differs from the pattern of Arr3a expression in the adult retina in which both red and green cones are labeled (Fig. 3P). This suggests that the expression of Arr3a in WT fish is delayed in green cones relative to red cones at 3 dpf, consistent with dim Arr3a expression in green cones relative to red cones previously observed at 4 dpf (31). Therefore, the lack of Arr3a expression in tdTomato⁺ cells in the *thrb*^{-/-} retina at 3 dpf suggests tdTomato⁺ precursors are transfated into a nonred cone (Arr3a⁻) photoreceptor subtype and that the

remaining Arr3a⁺ cells represent green cones in which Arr3a expression appears to be accelerated (Fig. 6D). Overall, our results in Fig. 6B and D suggest that in the absence of *thrb*, tdTomato⁺ precursors are transfated into a nonred/nongreen photoreceptor subtype and into cells that resemble horizontal cells.

To evaluate the possibility that tdTomato⁺ precursors in the *thrb*^{-/-} retina are transfated into UV cones, we used antibody staining to compare the number of UV cone opsin⁺ cells in WT and *thrb*^{-/-} retinas at 3 dpf. We found a significant increase in the number of UV cone opsin⁺ cells in *thrb*^{-/-} retinas compared to WT (Fig. 6F). Application of the UV cone opsin antibody to WT and *thrb*^{-/-} fish carrying the *trp2:tdTomato* transgene confirmed that a large subset of *trp2:tdTomato*⁺ precursors are transfated into UV cone opsin⁺ cells (closed arrow in Fig. 6E and G). We also observed a second population of UV cone opsin⁺ cells in the *thrb*^{-/-} retina which are tdTomato⁻; these cells likely represent native UV cones (open arrow in Fig. 6E). In further confirmation of these results, we observed that nearly all *trp2:tdTomato*⁺ cells in the ONL of the *thrb*^{-/-} retina are positive for the *sws1:GFP* transgene which exclusively marks UV cones (Fig. 6I–K) (32). The latter finding suggests that *trp2:tdTomato*⁺ cells in the *thrb*^{-/-} retina are exclusively converted into UV cones and not other photoreceptor subtypes within the ONL.

In addition to these findings, we observed that ~32% (mean \pm 0.04 SD, $n = 5$) of tdTomato⁺ cells in the larval *thrb*^{-/-} retina reside in the INL (asterisk in Fig. 6B). In contrast, we never observed tdTomato⁺ cells in the INL of WT retina (Fig. 6H). The localization of these tdTomato⁺ cells at the outer edge of the INL suggested that they might be horizontal cells. Indeed, we found that tdTomato expression in the INL colocalizes with expression of the *ptfla:GFP* transgene, a marker of horizontal cells (Fig. 6L–N) (30). These cells may represent new horizontal cells derived from cells originally fated to be red cones. Alternatively, it is possible that these cells are normal horizontal cells with aberrantly perdurant tdTomato expression.

In order to determine if transfating of *trp2:tdTomato*⁺ red cone precursors to UV cones also occurs in the growing retina of mature *thrb*^{-/-} fish, we analyzed coexpression of the UV cone-specific *sws1:GFP*⁺ transgene and *trp2:tdTomato*⁺ near the ciliary marginal zone (CMZ) of 12 dpf zebrafish. We find that multiple *thrb:tdTomato*⁺ cells colocalize with *sws1:GFP* near the *thrb*^{-/-} CMZ, suggesting that red cone precursors are transfated to UV cones when new retina is added during zebrafish growth (Fig. 6O–U). Despite the presence of these double-positive cells, we observed no evidence of supernumerary UV cones within the cone mosaic of adult *thrb*^{-/-} fish (Fig. 3K and O). We speculate

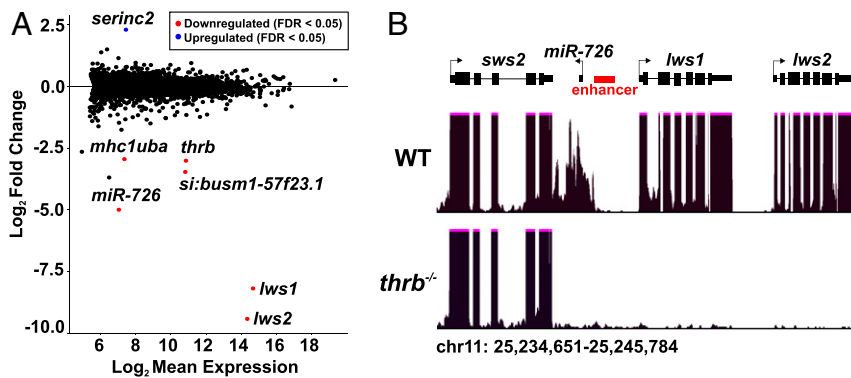


Fig. 5. Expression of only a very small number of genes is dysregulated in adult *thrb*^{-/-} retina. (A) Plot comparing gene expression in 6-mo-old WT and *thrb*^{-/-} retina obtained via RNA-seq (WT $n = 3$, *thrb*^{-/-} $n = 3$). Only five genes (excluding *thrb*) are significantly down-regulated in the *thrb*^{-/-} retina. *miR-726*, *lws1*, and *lws2* are previously characterized red-cone-specific genes. *mhc1uba* and *si:busm1-57f23.1* are putative red cone-enriched genes. (B) Screenshots of University of California Santa Cruz genome browser tracks demonstrate a selective loss of transcripts corresponding to *lws1*, *lws2*, and *miR-726*, and a normal number of transcripts corresponding to blue cone opsin (*sws2*). The enhancer labeled in red was previously shown to coordinately regulate *lws1* and *lws2* in zebrafish (4). In medaka, the orthologous enhancer regulates *miR-726* and *LWS-A*, the single medaka ortholog of *lws1/2* (27). *thrb*^{-/-} = *thrb*^{st1628/st1628}. FDR, false discovery rate (adjusted *P* value).

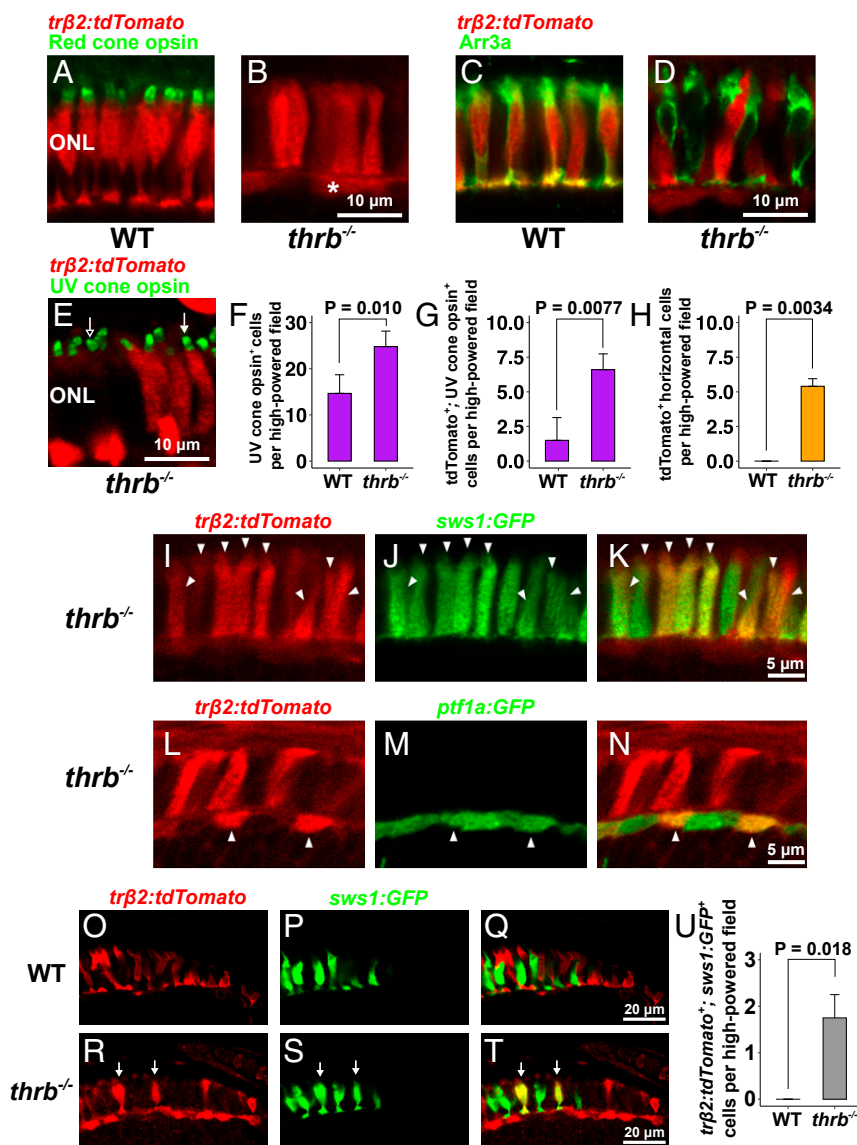


Fig. 6. Red cone precursors are transdifferentiated into UV cones in the larval and CMZ-derived *thrb*^{-/-} retina. (A–D) Optical sections of the ONL of 3 dpf *trβ2:tdTomato*⁺ and *trβ2:tdTomato*⁺;*thrb*^{-/-} larvae. (A and B) In the *thrb*^{-/-} retina, *trβ2:tdTomato*⁺ cells that reside in the ONL lack red cone opsin expression. Those cells that reside in the outermost INL resemble developing horizontal cells (asterisk). (C and D) In the *thrb*^{-/-} retina, *trβ2:tdTomato*⁺ cells do not express *Arr3a* ($n = 4$ WT, $n = 5$ *thrb*^{-/-}). (E) Confocal projection image of the ONL of 3 dpf *trβ2:tdTomato*⁺;*thrb*^{-/-} larvae shows that the majority of *trβ2:tdTomato*⁺ cells in the ONL are UV cone opsin⁺ (closed arrow). UV cone opsin⁺ cells that are *trβ2:tdTomato*⁻ (open arrow) are likely native UV cones. (F–H) Quantification of cell populations in the 3 dpf WT and *thrb*^{-/-} retina across 76 μm (mean \pm SD, $n = 6$ WT, $n = 5$ *thrb*^{-/-}). (F) The number of UV cones is increased in the *thrb*^{-/-} retina. (G) The number of *trβ2:tdTomato*⁺; UV cone opsin⁺ cells is increased in the *thrb*^{-/-} retina, suggesting that *trβ2:tdTomato*⁺ red cone precursors are transdifferentiated into UV cones in the *thrb*^{-/-} background. (H) The number of *trβ2:tdTomato*⁺ cells in the outer part of the INL (confirmed to be *ptf1a:GFP*⁺ horizontal cells in L–N) is increased in *thrb*^{-/-} retinas, suggesting that a subset of *trβ2:tdTomato*⁺ red cone precursors have been transdifferentiated into horizontal cells. (I–N) Optical sections of the retina of 3 dpf *trβ2:tdTomato*⁺;*thrb*^{-/-} larvae. (I–K) Almost all *trβ2:tdTomato*⁺ cells that reside in the ONL colocalize with *sws1:GFP* (arrowheads) and are therefore transdifferentiated into UV cones ($n = 3$ retinas). (L–N) *trβ2:tdTomato*⁺ cells in the INL colocalize with *ptf1a:GFP* (arrowheads), suggesting that they may represent transdifferentiated horizontal cells. (O–U) Red cone precursors are transdifferentiated into UV cones in the CMZ-derived *thrb*^{-/-} retina (O–T). Optical sections of the ONL immediately adjacent to the temporal CMZ of 12 dpf *trβ2:tdTomato*⁺ and *trβ2:tdTomato*⁺;*thrb*^{-/-} retina, tdTomato stained with anti-tdTomato antibody (O–Q). In the WT retina, the *sws1:GFP* transgene is not expressed in *trβ2:tdTomato*⁺ cells (R–T). In the *thrb*^{-/-} retina, a subset of *trβ2:tdTomato*⁺ cells are *sws1:GFP*⁺ (arrows), suggesting red cone precursors are transdifferentiated into *sws1:GFP*⁺ UV cones. The *sws1:GFP*⁺ cells that are *trβ2:tdTomato*⁻ are likely native UV cones. (U) The number of *trβ2:tdTomato*⁺; *sws1:GFP*⁺ cells across 94 μm immediately adjacent to the CMZ is increased in the 12 dpf *thrb*^{-/-} retina (mean \pm SD, $n = 4$ /group). All statistical tests were performed using Mann–Whitney U test. *thrb*^{-/-} = *thrb*^{st1627/st1627}.

that transdifferentiated UV cones derived from adult CMZ die prior to formation of the adult mosaic.

Discussion

TH nuclear receptors play a crucial role in at least two aspects of long-wavelength vision in zebrafish: determination of red cone

fate and control of *cyp27c1* expression. Here, we show that in the retina, *thrb* regulates the fate of red cone precursors, which in the absence of *thrb* transdifferentiate into UV cones and horizontal cells. We also show that in the RPE, TH acts through all three TH receptors (*thraa*, *thrab*, and *thrb*) to regulate *cyp27c1* expression and the resulting production of vitamin A₂-based retinoids. Recent

studies indicate that TH signaling also enhances expression of red-shifted paralogs of green and red cone opsin in both salmonids and zebrafish (14, 15). Thus, TH signaling coordinates multiple aspects of long-wavelength vision.

Comparison of the transcriptomes of WT and *thrb*^{-/-} retinas revealed only five down-regulated genes, suggesting the existence of very few red cone-specific genes. This finding contrasts with the discovery of hundreds of differentially expressed genes between cone subtypes in developing chicken retina (33). The greater degree of transcriptomic divergence between avian cone subtypes might be attributable to the greater number of structural differences between subtypes, including the presence/absence of intracellular organelles such as oil droplets and paraboloids (34, 35). Additionally, avian cone subtypes show differences in oil droplet pigmentation which is mediated by the differential expression of carotenoid-metabolizing enzymes and transporters (34, 36, 37). An alternative interpretation of our findings is that analysis of whole retina (rather than sorted cone subtypes) may have limited our ability to detect red cone-specific genes. Clearly, further studies will be required to determine the full extent of transcriptome divergence between zebrafish cone subtypes.

The transfecting of red cone precursors to UV cones in the *thrb*^{-/-} zebrafish retina suggests that *thrb* may have a similarly conserved developmental role in other vertebrate species. Studies in mouse have shown that in the *Thrb2*^{-/-} retina, *Opn1mw* (orthologous to zebrafish *lws1/2*) expression is reduced and *Opn1sw* (orthologous to zebrafish *sws1*) expression is increased (17). These results could be explained in part by a transfecting event similar to that observed in *thrb*^{-/-} zebrafish retina. Although we observe transfecting to UV cones in both embryo-derived and CMZ-derived *thrb*^{-/-} retina, supernumerary UV cones are not detected in the definitive adult mosaic, suggesting that transfecting UV cones die soon after birth. By comparing the transcriptome of native and transfecting UV cones, we might discover novel factors required for the viability of native UV cones.

TH has long been known as a potent physiological regulator. For example, it plays a major role in amphibian metamorphosis and in the induction of physiological changes associated with the parr-to-smolt transition of migratory salmon (38–40). Many of these TH-induced changes serve to promote adaptation to new environments (38–40). Although nothing is known about how TH regulates the A₁-to-A₂ switch in wild zebrafish, our work suggests that TH receptors may regulate *cyp27c1* expression under natural conditions. For example, day length or water temperature may influence *cyp27c1* expression in wild zebrafish, as these environmental factors have been shown to regulate the A₁-to-A₂ switch in other aquatic vertebrates (7).

In summary, we have found that TH receptors regulate two aspects of long-wavelength vision via the control of cone development and chromophore usage. The present study expands our understanding of the transcriptional mechanisms underlying these processes and suggests a coordinated mechanism by which TH facilitates visual system adaptation to red-shifted photic environments.

Materials and Methods

Mutant Zebrafish and Transgenic Lines. All procedures were carried out in accordance with the animal protocol approved by the Animal Studies Committee of Washington University. The *albino* line (*alb*^{b4/b4}) was used for ISH experiments (20). The *Tg(trj2:tdTomato)* transgenic line was a generous gift from Rachel Wong (University of Washington, Seattle, WA) (16). The *Tg(ptf1a:GFP)* transgenic line was a generous gift from Ryan Anderson (Indiana University School of Medicine, Indianapolis, IN) (30). The *Tg(sws1:GFP)* transgenic line, *Tg(zfsws1-5.5A:EGFP)^{tkk002}*, was acquired from

the Zebrafish National BioResource Project (<https://shigen.nig.ac.jp/zebra/>) (32). The *thraa*^{vp33rc1} and *thrab*^{vp31rc1} lines have been previously described in ref. 22. The *thrb*^{st1627} and *thrb*^{st1628} alleles were generated in-house using CRISPR-Cas9 with gRNAs targeting exon 9 of *thrb*. Further details are described in *SI Appendix, Materials and Methods*.

In Situ Hybridization. Anti-sense probes were prepared for *thraa*, *thrab*, *thrb*, and *rpe65a*. Detailed information is provided in *SI Appendix, Materials and Methods*.

Quantitative RT-PCR. WT and mutant fish were treated with TH (750 μL 400 μg/mL L-thyroxine in 0.1 M NaOH per liter of water for a final concentration of 300 μg/L) or vehicle control (750 μL of 0.1 M NaOH per liter of water), and *cyp27c1* expression was quantified using RNA extracted from eyes after removal of the lens. The expression of *thraa*, *thrab*, and *thrb* was quantified using RNA extracted from dissected retina and RPE. Further details are described in *SI Appendix, Materials and Methods*.

Antibodies and Immunohistochemistry. Primary antibodies used were as follows: rabbit polyclonal anti-DsRed (1:200; 632496, TakaraBio), which recognizes tdTomato (16); mouse monoclonal zpr1 (1:50; Zebrafish International Resource Center), which recognizes Arr3a (41, 42); rat monoclonal 10C9.1 (1:50, gift from Ted Allison, University of Alberta, Edmonton, Alberta, Canada), which recognizes UV cone opsin (43); mouse monoclonal 1D4 (1:50, AB5417, Abcam), which recognizes red cone opsin (44); chicken polyclonal anti-Arr3b (24) (1:250; gift from Stephan Neuhaus, University of Zurich, Zurich, Switzerland); and rabbit polyclonal anti-green cone opsin (45) (1:500, gift from David Hyde, University of Notre Dame, Notre Dame, IN). Secondary antibodies were as follows: donkey polyclonal anti-rat IgG conjugated with Alexa-488 (1:200, A21208, Invitrogen); goat polyclonal anti-mouse IgG conjugated with Alexa-488 (1:200, A11029, Invitrogen); goat polyclonal anti-chicken IgY conjugated with Alexa-488 (1:200; A11039, Invitrogen); donkey polyclonal anti-rabbit IgG conjugated with Alexa-488 (1:500; A21206, Invitrogen); donkey polyclonal anti-rabbit IgG conjugated with Alexa-555 (1:1,000; A31572, Invitrogen); and donkey polyclonal anti-mouse IgG antibody conjugated with Alexa-555 (1:200; A31570, Invitrogen). Further details on immunostaining are described in *SI Appendix, Materials and Methods*.

High Performance Liquid Chromatography. Whole eyes from vehicle and TH-treated WT and mutant zebrafish were collected and homogenized. Retinaldehydes were then derivatized to oximes using hydroxylamine treatment (Sigma, 255580). The quantities of A₁- and A₂-based retinaldehydes were then calculated based on all-*trans*-retinaldehyde oxime standards. Additional details are provided in *SI Appendix, Materials and Methods*.

RNA-Seq. For RNA-seq, 6 mo old WT and clutch-matched *thrb*^{st1628/st1628} zebrafish retina were isolated. Two retinas from a male and two retinas from a female zebrafish were combined to make one sample. Three WT and *thrb*^{st1628/st1628} samples were collected and RNA was extracted. Further details regarding RNA extraction, library preparation, sequencing, and data analysis are provided in *SI Appendix, Materials and Methods*.

Data Availability. RNA-seq data are available under GEO accession number GSE143312 (46). Analysis of differentially expressed genes in adult WT and *thrb*^{st1628/st1628} retina is available in [Dataset S1](#).

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