**Developmental Neuroscience**

**Foxq2 determines blue cone identity in zebrafish**

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Most vertebrate lineages retain a tetrachromatic visual system, which is supported by a functional combination of spectrally distinct multiple cone photoreceptors, ultraviolet (UV), blue, green, and red cones. The blue cone identity is ensured by selective expression of blue (sws2) opsin, and the mechanism is poorly understood because sws2 gene has been lost in mammalian species such as mouse, whose visual system has been extensively studied. Here, we pursued loss-of-function studies on transcription factors expressed predominantly in zebrafish cone photoreceptors and identified Foxq2 as a blue cone–specific factor driving sws2 gene expression. Foxq2 has dual functions acting as an activator of sws2 transcription and as a suppressor of UV (sws1) opsin transcription in blue cones. A wide range of vertebrate species retain both fox2 and sws2 genes. We propose that Foxq2-dependent sws2 expression is a prevalent regulatory mechanism that was acquired at the early stage of vertebrate evolution.

**INTRODUCTION**

Most vertebrates have highly developed camera-type eyes with duplex retinas equipped with rod and cone photoreceptor cells (1–3). Rods with a higher light-sensitivity respond to single photons and mediate scotopic vision under twilight conditions at night. In contrast, cones show a relatively lower sensitivity without saturating in brighter light and mediate photopic vision under a daylight condition. Color discrimination is established by a combination of spectrally distinct cone subtypes, each expressing a single cone opsin out of four subfamilies: ultraviolet-sensitive [SWS1, wavelength of maximum sensitivity (λmax): 360 to 420 nm], blue-sensitive (SWS2, λmax: 400 to 470 nm), green-sensitive (RH2, λmax: 460 to 510 nm), and red-sensitive opsins (LWS, λmax: 510 to 560 nm) (4, 5). Most vertebrates retain the tetrachromatic visual system organized by the four cone opsin subfamilies. A full set of genes encoding the four cone opsins is present in the Southern Hemisphere lamprey, a jawless vertebrate belonging to the earliest-branching vertebrate group (6). This fact supports the idea that the last common ancestor of vertebrates should have had color vision based on the four cone opsin subfamilies (4, 7).

Retinal progenitor cells differentiate into all types of retinal neurons in a temporal order, conserved among many species (8, 9). In the later process, transcription factors regulate photoreceptor-specific gene expression. Cone-rod homeobox (Crx) is an upstream transcriptional regulator for both rod and cone photoreceptors (10). A rod master regulator, neural retina leucine zipper (NRL), and its downstream factor, nuclear receptor 2E3 (Nr2e3), enhance rod-specific gene expression and repress sws1 expression (11, 12). With regard to cone subtypes, thyroid hormone receptor beta (Thrb) is a master transcriptional regulator for expression of lws opsin and responsible for differential expression between lws and sws opsins in mice (13), zebrafish (14), and human (15). Another transcription factor, T-box 2b (Tbx2b), plays an essential role in sws1 opsin expression in zebrafish (16). On the other hand, much less is known about a regulatory network governing expression of the middle-wavelength–sensitive opsin genes, sws2 and rh2, which have been lost in most mammalian species.

In zebrafish, a tetrachromatic freshwater fish, we found that sine oculis homeobox 7 (Six7) is required for expression of all the four subclasses of rh2 genes (rh2-1, rh2-2, rh2-3, and rh2-4) (17), which are tandemly arrayed, expressed in different cone cells, and spectrally distinct from each other (18, 19). Six7 and its homolog Six6b control sws2 expression as well (20). The cone-enriched transcription factors, Six7 and Six6b, share common DNA binding sites in both rh2 and sws2 gene loci, and Six6b overexpression rescues the reduced level of rh2 expression in six7 deficient fish (20). The overlapping functions between Six6b and Six7 for sws2 and rh2 expression imply that an additional factor(s) directs selective expression of either sws2 or rh2 genes and determines the identities of the middle-wavelength–sensitive cone subtypes. In the present study, gene expression profiling with isolated rods and cones enabled us to identify a list of cone-enriched transcription factors. Our in vivo functional analyses revealed a core transcriptional network, in which Foxq2 acts as a downstream regulator of Six7 and regulates sws2 expression. We demonstrate that Foxq2 is a terminal selector determining SWS2 cone identity during development of the middle-wavelength–sensitive cone subtypes.

**RESULTS**

A severe reduction of sws2 expression in foxq2 mutant zebrafish

Six6b and Six7 are predominantly expressed in zebrafish cone photoreceptors and responsible for expression of the middle-wavelength–sensitive opsin genes, sws2 (blue) and rh2 (green) (20). To identify a transcription factor(s) that governs differentiation between SWS2 and RH2 cone subtypes, we searched for cone-specific genes by comparing gene expression profiles between cones and rods. These photoreceptor cells were purified from the retinas of transgenic adult zebrafish, each of which express enhanced green fluorescent protein (EGFP) in all cone subtypes [Tg(gnat2:egfp)] or in rods [Tg(rh2:egfp)] (17). The cone or rod enrichment in these purified samples was validated by reverse transcription quantitative polymerase chain reaction (RT-qPCR) analyses of cone- and rod-specific transducin...
alpha-subunit genes, i.e., \textit{gnat2} and \textit{gnat1}, respectively (Fig. 1A) (17). A subsequent microarray analysis revealed approximately 500 genes showing more than 10-fold higher expression in cones than in rods (data file S1). These cone-enriched genes included four transcription factors, \textit{foxq2}, E2F transcription factor 7 (\textit{e2f7}), nuclear factor 1A (\textit{nfia}), and nuclear receptor 2F6b (\textit{nr2f6b}), the roles of which in photoreceptor development were not known. Genomic loci of these genes harbor Six6b- and Six7-binding sites as revealed by our chromatin immunoprecipitation sequencing analysis (fig. S1) (20), implying that some of these cone-enriched transcription factors mediate a regulatory function(s) downstream of Six6b and Six7. In addition, we paid attention to two cone-enriched transcription factors, \textit{tbx2b} and \textit{thrb}. They are known to be required for expression of \textit{sws1} (16) and \textit{lws} (14), respectively, but their contributions to the \textit{sws2} and \textit{rh2} gene expression have not been well characterized.

Cone-enriched expression of these six transcription factors was verified by RT-qPCR analyses with purified cone and rod cells (Fig. 1A). We generated loss-of-function mutants of zebrafish for each of the six transcription factors by introducing a frameshift mutation. In these mutants, ocular transcript levels of the middle-wavelength-sensitive opsin genes were examined by RT-qPCR analysis (Fig. 1B). Among the mutant larvae, a \textit{foxq2} mutant (Fig. 1C, ja74) displayed the most notable reduction in \textit{sws2} expression as compared to the wild-type (WT) siblings (Fig. 1B). Another mutant line of \textit{foxq2} (Fig. 1C, ja77) similarly showed a severe reduction of \textit{sws2} expression in the larvae (fig. S2A). The decrease of \textit{sws2} expression in the mutant larvae (Fig. 1D and fig. S2A) was not due to delayed development of SWS2 cones because the adult \textit{foxq2} mutant (ja77) also exhibited minimal expression of \textit{sws2} gene (fig. S2B). In the \textit{foxq2} mutant retinas, in situ hybridization signals of \textit{sws2} transcripts were

![Fig. 1. The loss-of-function analysis for cone-enriched transcription factors.](image-url)
undetectable with no apparent change in retinal morphology at the larval and adult stages (figs. S2C and S3A). These results demonstrate that foxq2 is indispensable for sws2 expression in SWS2 cone subtype.

The foxq2 mutants (ja74 and ja77) were not only deficient in sws2 expression but also characterized by significant reduction in mRNA level of arrestin 3b (arr3b), a cone phototransduction gene selectively expressed in SWS1 and SWS2 cone subtypes (Fig. 1D and fig. S2) (21). sws1- and arr3b-expressing cone cells in the larval eye were visualized by in situ hybridization chain reaction (HCR) (Fig. 2A). Almost all arr3b-positive cells in the foxq2 mutant coexpress sws1, suggesting the absence of arr3b-positive and sws1-negative cells, i.e., SWS2 cones, in the foxq2 mutant. We then assessed mosaic arrays of cone photoreceptor cells in the adult retina, in which a SWS2 cone (B in Fig. 2B) is flanked by RH2/LWS double cones (G/R) in each mosaic unit (G/R/B/R/G/V; see Fig. 2B and fig. S3B) (22). Immunohistochemistry and in situ HCR labeling in the flat-mounted retina revealed that SWS2 cone is selectively lost in each mosaic unit of the foxq2 mutant retina (G/R/B/R/G/V; Fig. 2, B and C, and fig. S3, B and C). The expression of cone-specific phototransduction gene, gnat2 (in SWS1/SWS2/RH2/LWS cones), was colocalized with mRNA expression of either sws1 (in SWS1 cones) or arr3a (in RH2/LWS cones) in the mutant. These observations support that the mutant retina contains SWS1, RH2, and LWS cones and has no “opsin-empty” cones (Fig. 2C and fig. S3C). In the foxq2 mutant retina, the number of apoptotic cells [terminal deoxynucleotidyl transferase–mediated deoxyuridine triphosphate nick end labeling (TUNEL)–positive cells] was increased in the outer nuclear layer of the central

Fig. 2. The loss of SWS2 cones in the foxq2 mutant. (A) Expression patterns of sws1 and arr3b (SWS1 and SWS2 cones) in 5-dpf larval eyes of the foxq2 mut (ja74) examined by in situ HCR. Magnified view (a box surrounded with white lines) is indicated in the right side of each panel. Scale bars, 20 μm. (B) Fluorescent images of the flat-mounted retinas prepared from the adult WT, the foxq2 mut (ja74), and Tg(-3.5opn1sw2:EGFP)kj11Tg (sws2:egfp), where EGFP is expressed in SWS2 cones (green). The retinas were immunostained with zpr1 antibody (arr3a, red) and also stained with DRAQ5 to highlight cell nuclei (blue). V, SWS1 cone; B, SWS2 cone; G, RH2 cone; R, LWS cone. Scale bar, 10 μm. See also fig. S3B. (C) Expression patterns of phototransduction genes in the flat-mounted retinas examined by in situ HCR. See also fig. S3C. (D) Left: Fluorescent images in retinal cryosections from the adult fish labeled for terminal deoxynucleotidyl transferase–mediated deoxyuridine triphosphate nick end labeling (TUNEL) (red). The cell nuclei were counterstained with 4′,6-diamidino-2-phenylindole (DAPI) (blue). Scale bar, 50 μm. Right: Quantification of TUNEL-positive cells in the central and peripheral retina. The numbers of TUNEL-positive cells were counted for each cryosection and averaged (means ± SEM, n = 80 for WT, n = 63 for the foxq2 mut; *P < 0.05, Student’s t test). See also fig. S3D. (E) Expression patterns of sws2, rh2-1, and rh2-2 in 5-dpf larval eyes of WT and foxq2 mut (ja74) examined by in situ HCR. The number of opsin gene–positive cells in the central region of the retina is indicated in a bar graph. The number in the upper-right corner for each panel represents the unique identity of the eye. Data are represented by means ± SEM (n = 3). Scale bar, 50 μm. See also fig. S3E.
region and in the ciliary marginal zone of the peripheral region (Fig. 2D and fig. S3D). These observations suggest that foxq2 deficiency impairs formation and/or maturation of SWS2 cones.

In parallel, the foxq2 mutation caused significant increase in mRNA levels of two major rh2 subclasses, rh2-1 in the larvae (fig. S2A, ja77) and rh2-2 in the adult (fig. S2B, ja77), as well as up-regulation of rh2-2 in the larvae (Fig. 1D, ja74, and fig. S2A, ja77) and rh2-4 in the adult (fig. S2B, ja77). In contrast, the numbers of rh2-1 and rh2-2–expressing cone cells in the foxq2 mutant larvae were similar to those in WT as revealed by in situ HCR analysis (Fig. 2E and fig. S3E).

The number of RH2 cones also seemed normal in the adult foxq2 mutant retina (Fig. 2, B and C), in which each cone mosaic unit contains two RH2/LWS double cones as observed in WT. In addition, we observed no substantial expression of rh2-1 and rh2-2 outside the photoreceptor layer in the mutant retina (figs. S2C and S3A). It is most probable that rh2 expression is up-regulated in the individual RH2 cones of the foxq2 mutant retina.

Tbx2b is also a cone-specific transcription factor (Fig. 1A) and is required for expression of sws1 gene (16). In the present study, tbx2b<sup>k20</sup> mutant was designed to encode C-terminally truncated Tbx2b protein (Fig. 1E), which lacked two DNA recognition helices of T-box domain (23) in a manner similar to tbx2b<sup>l57</sup> mutant (24). As reported in tbx2b<sup>b</sup> (24), tbx2b<sup>j20</sup> larvae exhibited a severe decrease of sws1 and a parallel increase of rhodopsin (rho) expression (Fig. 1F). Notably, we noticed that tbx2b<sup>k20</sup> mutation caused 40% reduction of sws2 expression level from that in the WT siblings and substantial increase in mRNA levels of rh2-1 and rh2-2 (Fig. 1, B and F). Although the effect of tbx2b mutation on ssw2 expression was weaker than that of foxq2 mutation, tbx2b appears to play a supportive role to foxq2 on ssw2 expression in addition to the essential role on ssw1 expression.

Thrβ is another cone-specific transcription factor (Fig. 1A), and its mutant thrb<sup>ja27</sup> (fig. S4A) manifested a massive reduction of both lws1 and lws2 expression and a concomitant increase in ssw1 expression in the larval eyes (Fig. 1B and fig. S4B) as reported in previous studies (14, 15, 25). We noticed that thrb<sup>ja27</sup> mutant exhibited a moderate but significant increase in ssw2 gene expression (Fig. 1B and fig. S4B). The ssw2 up-regulation accompanied no significant reduction of rh2 expression level (Fig. 1B and fig. S4B). Accordingly, Thrβ contributes to fine-tuning of ssw2 expression together with a dominant role in regulation of lws expression. In contrast, nfa, nrf2f6b, and e2f7 mutant fish showed no noticeable change in ssw2 or rh2 expression (Fig. 1B and fig. S4, C to H). These three transcription factors are dispensable for middle-wavelength cone opsin gene expression.

Collectively, our mutant analysis demonstrates that ssw2 expression is predominantly regulated by a cone-specific transcription factor, Foxq2, with the aid of regulation by Tbx2b and Thrβ. It is most probable that Foxq2 is also required for formation and/or maturation of SWS2 cones.

**SWS2 cone subtype–specific expression of foxq2**

To gain insights into how foxq2, tbx2b, and thrb regulate ssw2 and rh2 expression, we investigated gene expression profiles of foxq2, tbx2b, and thrb among the four cone subtypes: SWS1, SWS2, RH2, and LWS. These four cone cells were isolated from four different lines of transgenic fish, each of which expresses EGFP in one of the four cone subtypes (Fig. 3A; see Materials and Methods for details). The cone subtype enrichment in these purified samples was validated by RT-qPCR analyses of cone opsin genes (Fig. 3B). Subsequent analysis of foxq2, tbx2b, thrb, six6b, and six7 in the four samples revealed that foxq2 was specifically expressed in SWS2 cone subtype (Fig. 3C). Another SWS2 regulator, tbx2b, was expressed in both SWS1 and SWS2 cone subtypes, whereas thrb was expressed only in the LWS cone subtype (Fig. 3C). In contrast, SWS2/RH2 regulators, six6b and six7, were expressed in all the cone subtypes (Fig. 3C). The SWS2 cone subtype–enriched expression of foxq2 and tbx2b suggests that these two factors coordinate to regulate the cell type–specific expression of ssw2.

The expression of foxq2 gene in the larval stage was visualized by in situ HCR. The foxq2 transcript signals were colocalized with those of ssw2 (in SWS2 cones; Fig. 3D) but not with those of ssw1 (in SWS1 cones) or arr3a (in RH2/LWS cones) (Fig. 3E). foxq2 expression is thus specific to SWS2 cones among the cone subtypes at both larval (Fig. 3D) and adult (Fig. 3C) stages. RT-qPCR analysis revealed that foxq2 expression was selectively expressed in the retina among various adult tissues (Fig. 4A), which is consistent with the publicly available RNA sequencing data (fig. S5). In the larvae, foxq2 expression was detected only in the anterior half of the body (including the eyes) but not in the posterior half (Fig. 4B). The eye-specific foxq2 expression in the larvae was visualized by in situ HCR (Fig. 4C). Notably, the foxq2 signals were enriched in ciliary marginal zone (Fig. 4C), which contains a population of actively proliferating progenitor cells (26). This expression pattern suggests that foxq2 expression precedes ssw2 expression in differentiating cone cells. The mRNA level of foxq2 was severely reduced in six6a/six6b/six7 triple knockout (TKO) lacking ssw2 expression, i.e., loss of SWS2 cone identity from the retina (Fig. 4, D and E) (20). foxq2 expression was also abrogated in the larval eyes of foxq2 mutant (ja74) as compared to that of WT sibling (Fig. 4F). The close correlation between foxq2 and ssw2 expressions implies that Foxq2 is responsible for establishing SWS2 cone identity. Meanwhile, tbx2b was expressed widely in various tissues at the larval (Fig. 4B) and adult stage (Fig. 4A and fig. S5), and the ocular tbx2b expression level was unaffected in the foxq2 mutant (Fig. 4F) and six6a/six6b/six7 TKO (Fig. 4, D and E), both of which were deficient in ssw2 expression. These observations indicate dominant expression of tbx2b outside of SWS2 cones and imply that Tbx2b may have pleiotropic roles for eye development (27) including the cone identity determination.

**Functional interaction of Foxq2 with ssw2 promoter**

We then investigated functional interaction of Foxq2 with ssw2 promoter. Foxq2 protein (Fig. 1C) has a conserved DNA binding domain, termed forkhead domain composed of about 100 amino acid residues (28). In Fox transcription factor family, Foxq2 is categorized into clade I forkhead proteins (29), which recognize two types of common forkhead-target DNA motifs termed FkhP (RYAAAYA) and FkhS (AHAAAA) (30). Our motif scanning analysis (see Materials and Methods) revealed two FkhP motifs and two FkhS motifs present within 1.56-kb ssw2 promoter region (Fig. 5A and fig. S6), which drives selective gene expression in SWS2 cones (31). In a cell-based reporter assay, the 1.56-kb ssw2 promoter was transactivated by VP64-Foxq2 (Fig. 5, B and C), in which Foxq2 is N-terminally fused with four repeats of the VP16 transcriptional activator domain (Fig. 5B) (32). The transcriptional activation was attenuated by deletion of the upstream region of 1.03 or 1.26 kilo–base pair (kb) (0.53- or 0.3-kb ssw2 promoter, respectively), leaving single FkhS motif (Fig. 5, A and C). Still, we observed more than 30-fold
transactivation of the 0.3-kb sws2 promoter by VP64-Foxq2, while the activation was largely reduced by the complete deletion of the promoter sequence (Fig. 5C). The VP64-Foxq2–dependent transactivation of a shorter promoter (0.25-kb sws2 promoter) was markedly decreased by introducing a 6-bp mutation in the FkhS motif (Fig. 5D). We then generated a mutant protein, termed Fork-del, lacking nine amino acid residues that are required for specific DNA binding of the forkhead domain (Fig. 5B) (30). The protein mutation abolished the ability of the transcriptional activation of the sws2 promoter (Fig. 5, B, D, and E). These results demonstrated that Foxq2 functionally interacted with the forkhead-target DNA motif in the sws2 promoter. Together with the SWS2 cone–specific expression of foxq2 (Fig. 2), Foxq2-mediated transcriptional regulation of sws2 gene would reasonably account for the selective expression of sws2 opsin in SWS2 cone subtype.

**sws2 regulation by Foxq2 downstream of Six7**

To gain deeper insight into the mechanism underlying the determination of the SWS2 subtype identity, we generated foxq2 transgenic (foxq2-tg) zebrafish, Tg(-5.2crx:EGFP-2A-FLAG-foxq2), in which both EGFP and Foxq2 expressions are driven by 5.2-kb crx promoter in all the developing and matured photoreceptor cells (Fig. 6, A and B) (14). RT-qPCR analysis of larval ocular mRNAs in three lines of foxq2-tg (ja78Tg, ja79Tg, and ja91Tg) revealed that the forced expression of foxq2 in WT background severely reduced the sws1 mRNA level (Fig. 6C and fig. S7, A and B). This phenotype was not accompanied by any detectable change in the expression levels of tbx2b and crx (Fig. 6D and fig. S7, C and D), which are regulators of sws1 expression (Fig. 1K) (16). It is most probable that Foxq2 suppresses sws1 expression in cones. In in situ HCR analysis, the forced foxq2 expression largely decreased the number of SWS1 cones and increased that of SWS2 cones (Fig. 6E and fig. S7E), while the numbers of rh2-1– and rh2-2–expressing cells were similar between the foxq2-tg (ja78Tg) and WT (fig. S7F). It should be noted that the foxq2-tg retina contained a considerable number of SWS1/SWS2 hybrid cones (sws1/sws2 double-positive cells) (Fig. 6F and fig. S7E), whereas neither LWS/SWS2 hybrid nor RH2/SWS2 hybrid cones were observed in the foxq2-tg retina (Fig. 6F and fig. S7G). These results indicate that Foxq2 controls SWS2 subtype identity by promoting sws2 expression and suppressing sws1 expression (Fig. 6G).

The selective expression of foxq2 in SWS2 cones (Fig. 3, C and D), together with binding of Six6b and Six7 to foxq2 gene locus (fig. S1 and 20), suggests that sws2 expression is regulated by Foxq2 downstream of Six6b and Six7 (Fig. 6G). This possibility was explored by overexpressing foxq2 in the homozygous six7 KO (17), in which expression levels of sws2 and foxq2 are markedly reduced (Fig. 6, C and D). The forced expression of foxq2 in the six7 KO background (foxq2-tg;six7 KO) recovered the reduced expression of sws2 up to a level
higher than that in the WT control (Fig. 6C). In contrast, the over-expression of foxq2 had no significant effect on the severely reduced expression levels of rh2 (rh2-1 plus rh2-2) (Fig. 6C) or on mRNA levels of crx, tbx2b, six6b, and six7, transcriptional regulators of cone opsin expression (Fig. 6D). These results demonstrated that Foxq2 regulates sws2 expression downstream of Six7, suggesting that Foxq2 is a terminal selector for SWS2 subtype identity (Fig. 6G).

**Conservation of FOXQ2 gene among vertebrate species**

FOXQ2 gene is annotated in the genomes of a wide range of vertebrate species (Fig. 7), such as ray-finned fish (spotted gar, zebrafish, and medaka), a lobe-finned fish (coelacanth), and an avian (sparrowhawk), all of which retain SWS2 gene. We found highly conserved gene synteny around FOXQ2 locus among the vertebrates, and the synteny analysis of the human genome indicated the absence of any gene homologous to FOXQ2 harboring the forkhead domain highly conserved among FOXQ2 subfamily members (Fig. 7B). BLAST search and subsequent manual annotation revealed that a mammalian species, platypus, retains a gene orthologous to FOXQ2 subfamily members (Fig. 7 and fig. S8; see also Supplementary Text). The oviparous mammals including platypus diverged from marsupial and placental mammals at the earlier stage of mammalian evolution (around 200 million years ago). Notably, platypus retains SWS2 gene in its genome (33), whereas many other mammals have lost it (7, 34) most likely due to a long evolutionary history of nocturnality in mammalian ancestors (35). These lines of genomic evidence suggest that Foxq2-dependent SWS2 expression is a highly conserved regulatory mechanism that was acquired at the early stage of vertebrate evolution.

**DISCUSSION**

Under the solar light irradiation, both terrestrial and aquatic environments on the earth’s surface are enriched with blue-to-green region of visible spectrum (36), which is detected by cone photoreceptor cells expressing middle-wavelength–sensitive opsin genes, SWS2 and RH2. Their expression requires transcription factors, Six6 and Six7, in zebrafish (17, 20), and the six6/six7 mutant fish lacking both SWS2 and RH2 cones show severely reduced survival rate due to impairment of visually driven foraging behavior (20). A recent study of in vivo calcium imaging of zebrafish cone photoreceptors reported that SWS2 and RH2 cones, but not SWS1 or LWS cones, display strong spectral opponency and efficiently extract chromatic information from the natural light spectrum (37). In this way, a combination of spectrally distinctive SWS2 and RH2 cones particularly plays an important role in the tetrachromatic visual system. The present study explored the transcriptional regulatory logic that defines SWS2 and RH2 cone identities. For this purpose, cone-enriched transcription factors were comprehensively identified by the transcriptome analysis with purified cone cells (versus purified rod cells; data file S1). Subsequent functional analyses of these transcription factors demonstrated that Foxq2 is indispensable for sws2 expression and probable for formation and/or maturation of SWS2 cones (Fig. 1). We pursued expression profiling of the
cone-enriched transcription factors among the four isolated cone subtypes (Fig. 3, A to C). foxq2 is selectively expressed in SWS2 cone, whereas all cone subtypes express sixtb and six7 (Fig. 3C), which are required for both sws2 and rh2 gene expression (20). A transcriptional network is deciphered in which Foxq2 acts as a downstream regulator of Six7 (Fig. 6, A to D) and regulates sws2 transcription and as a suppressor of sws1 genes in SWS2 cones (Fig. 6C and fig. S2). Thus, Foxq2 has dual functions acting as an activator of sws2 transcription and as a suppressor of sws1 and rh2 genes in SWS2 cones (Fig. 6G) and in developing cones. The dual functions of Foxq2 would enhance robustness of SWS2 cone identity. Meanwhile, thrb is predominantly expressed in LWS cone subtype (Fig. 3C), being consistent with the widely accepted idea that Thrb is a master regulator for the LWS cone (13–15). Collectively, Foxq2, Tbx2b, and Thrb can be defined as the terminal selectors (38) that govern cell fate determination of SWS2, SWS1, and LWS cone subtypes, respectively.

In contrast to these three terminal transcriptional selectors, six7 is widely expressed among the four cone subtypes (Fig. 3C) and thus six7 is unlikely to be a terminal selector of RH2. Although Six7 is indispensable for expression of rh2 [rh2-1, rh2-2, rh2-3, and rh2-4; Fig. 6C and (17)], terminal differentiation of RH2 cone might be mediated by an unidentified transcription factor. An alternative idea is that Rh2 cone identity is established by both (i) the presence of six7 and (ii) the absence of the terminal selectors such as tbx2b (for SWS1), foxq2 (for SWS2), and thrb (for LWS). In this scenario, differentiation toward Rh2 cones should be a default pathway in the cone development because Rh2 cone differentiation is governed by six7 (17) and six7 expression begins as early as that of crx, a master
regulator of cone and rod development. It is worth noting that the "Rh2-default" hypothesis is consistent with molecular phylogeny of cone opsins genes, in which RH2 subfamily diverged from SWS2 at the latest step of the molecular evolution of the cone opsins. Building on these assumptions, we propose that the tetrachromatic color vision system in ancestral vertebrates used the RH2-default mechanism of cone differentiation as is observed in zebrafish. In mammals, on the other hand, RH2 and SWS2 genes have been lost, and hence, it is likely that the RH2-default mechanism has been modified to a "SWS1-default" mechanism.
latter mechanism, the presence or absence of only one terminal selector, THRB, directs differentiation between LWS and SWS1 cones as shown in mice (8, 13).

FOXQ2 gene is evolutionarily conserved not only among vertebrates but also in many invertebrates (29, 30, 41). In embryos of the invertebrates, FOXQ2 is responsible for specification and positioning of anterior neuroectoderm (42, 43). The anterior neuroectoderm develops into central nervous system, in which a subset of sensory cells express a photoreceptive protein, c-opsin, that is orthologous to vertebrate cone and rod opsins (44, 45). FOXQ2 might regulate differentiation of the c-opsin–expressing cells from progenitor cells in the anterior neuroectoderm. A transcriptional network in the anterior neuroectoderm of invertebrates also includes six3, a gene orthologous to vertebrate six3, six6, and six7 (42, 43). The presence of the Six3/6/7-Foxq2 transcriptional network in both vertebrate and invertebrate species suggests that their common ancestor used this transcriptional network for development of the light-sensitive cells. In the vertebrate lineage, the Six3/6/7-Foxq2 transcriptional network might have been co-opted for establishment of SWS2 cone identity during or after the appearance of a full set of cone opsins, thereby conferring high-acuity discrimination of blue-to-green region of visible light.

MATERIALS AND METHODS
Zebrasfish
The Ekkwill strain was used as the WT zebrafish. All appropriate ethical approval and licenses were obtained from Institutional Animal Care and Use Committees of The University of Tokyo. All the procedures were conducted according to the local guidelines of The University of Tokyo. Zebrafish were raised in a 14-hour light/10-hour dark cycle and fed twice per day with live baby brine shrimps. Embryos were raised at 28.5°C in egg water (artificial seawater diluted 1.5:1000 in water). Mutant strains and transgenic lines used in the present study are listed in table S1.

Purification of rod and cone photoreceptor cells
For isolation of rod and cone photoreceptor cells, we used the transgenic zebrafish lines, Tg(rho:egfp)ja2 (46) and Tg(gnata2:egfp)ja23 (17), which express EGFP in rods and all cone subtypes, respectively. The rod and cone cells were isolated as described previously (17, 20, 47). Briefly, retinas were dissected from dark-adapted adult fish under dim red light. The isolated retinas were digested with 0.25% trypsin, deoxyribonuclease I (10 U/ml), 2 mM MgCl2, and 2 mM EGTA in Ca2+-free Ringer’s solution for 30 min at 37°C. The reaction was terminated by adding soybean trypsin inhibitor (final 0.5%) and fetal bovine serum (final 10%). The dissociated cells were filtrated through a 35-μm nylon mesh (Falcon). EGFP-positive cells were isolated with a fluorescence activating cell sorter (FACSAria, BD Biosciences) by the following three parameters: forward scatter, side scatter, and green fluorescence. The isolated cells were directly collected into 800 μl of TRIzol reagent (Thermo Fisher Scientific) in 1.5-ml microtubes.

For isolation of cone subtypes, we used four lines of transgenic zebrafish: Tg(-5.5opn1sw1:EGFP)ja9 (48), Tg(3.5opn1sw2:EGFP)ja11 (31), Tg(RH2-2/GFP-PAC)ja4 (49), and Tg(-0.6opn1lw1-lws2:GFP)ja19.
Microarray
Total RNA was isolated from the sorted cells using an RNeasy Extraction kit (Qiagen). Quality and quantity of the resulting RNA were assessed using a NanoDrop ND-2000 spectrophotometer (Thermo Fisher Scientific) and an Agilent 2100 Bioanalyzer (Agilent Technologies). Microarray analysis was performed using the Agilent 4×44 k Zebrafish microarray according to the manufacturer’s protocol for the two-color method. Cy3- or Cy5-labeled complementary RNA (cRNA) probe was synthesized from 150 ng of total RNA using the Quick-Amp Labeling Kit (Agilent Technologies). Quantity of the resulting labeled cRNA was assessed using a NanoDrop ND-2000 spectrophotometer. Equal amounts of Cy3- and Cy5-labeled cRNA (825 ng) from two different samples were hybridized to zebrafish microarrays (Agilent Zebrafish Oligo Microarrays version 2, G2519F-019161) for 17 hours at 60°C. To compare gene expression profiles between cone and rod samples, we analyzed two independent biological replicates as follows: (i) Cy3-rod#1 cRNA and Cy5-cone#1 cRNA and (ii) Cy5-rod#2 cRNA and Cy3-cone#2 cRNA. The hybridized arrays were then washed and scanned using an Agilent microarray scanner (G2505 C; Agilent Technologies). Data were extracted from the scanned image using Feature Extraction version 10.5.1.1 (Agilent Technologies). We then listed differentially expressed genes with Microsoft Excel as follows: (i) We excluded any probes whose signals in two arrays were all determined as negative, meaning that probes were not flagged as “WellAboveBG” in any of two arrays. (ii) We selected all the probes whose signal intensities vary largely between the photoreceptor cell types according to the following thresholds of the averaged ratios: 10-fold increase for cone compared with rod and 4-fold increase for rod compared with cone. (iii) We checked whether both of the two independent biological replicates showed similar expression profiles and selected probes whose absolute values of the ratios in the two biological replicates were both >2.0 for rod/cone or both >3.0 for cone/rod. The lower threshold was given for the rod/cone ratio because a small but noticeable level of rod contamination was detected in cone samples such as gnat1 (Fig. 1A). We set these threshold values on the basis of the ratios observed for known cone-specific genes. The microarray datasets are available at the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database (GEO no. GSE168749).

RT-qPCR analysis
Zebrafish were anesthetized by chilling on ice, and their tissues were collected during the light phase of the light-dark cycle and soaked in RNAlater (Sigma-Aldrich) for RT-qPCR analysis or fixed with 4% paraformaldehyde (PFA) in Ca2+- and Mg2+-free Dulbecco’s phosphate-buffered saline (D-PBS) for cryosectioning.

Generation of mutant zebrafish
The foxq2, thrb, and nrz2f6b mutants were generated with a CRISPR-Cas9 system. The single-guide RNA (sgRNA) sequences (Table S2) were designed to target exon 1 or exon 2 using CRISPRdirect (https://crispr.dbcls.jp/), CHOPCHOP (https://chopchop.cbu.ubc.no), or CRISPRscan (www.crisprscan.org). The sgRNA was synthesized by a cloning-free method as previously described (51). For generating Cas9 mRNA, the template plasmid DNA, pCS2–Cas9n (Addgene, no. 47929) (52) or pCS2+hSpCas9 (Addgene, no. 51815) (53), was used for in vitro transcription. Cas9 mRNA was synthesized with the SP6 mMESSAGE mMACHINE Kit (Ambion) and purified with the RNeasy Mini kit (Qiagen). The solution containing 200-pg Cas9 mRNA and 25-pg sgRNA was injected into the cytoplasm of the one cell-stage embryos.

The tbx2b, e2f7, and nfia mutants were generated by transcription activator–like effector nucleases (TALENs) as previously described (17, 20). To target each of the three genes, a pair of the TALE effector repeats (Table S2) were designed to target exon 2 or exon 3 with the Golden Gate assembly methods. TALEN mRNA was synthesized using the SP6 mMESSAGE mMACHINE Kit (Ambion) and purified with the RNeasy Mini kit (Qiagen). The solution containing 200 pg each of the two TALEN mRNAs was injected into the cytoplasm of the one-cell stage embryos.

The injected fish (F0) were crossed with the WT zebrafish. The resultant F1 offspring were screened for the presence of CRISPR-Cas9– or TALEN-induced mutations by a combination of PCR and subsequent enzyme digestion for thrb, nrz2f6b, tbx2b, e2f7, and nfia mutants or by heteroduplex mobility assay for foxq2 mutants as described previously (54). To sequence-verify mutations, genomic sequences surrounding the mutations were amplified by nested PCR, and the resultant PCR products were sequenced directly. After isolation of mutant zebrafish and verification of the mutations, the mutant genotypes were confirmed by a combination of PCR and subsequent enzyme digestion except for the foxq2 (ja74) gene locus. The foxq2ja74 mutant genotype was determined by PCR with two pairs of primers: (i) foxq2_Fw1A (5’-TGGCT AACAT ACAG AACAA CAACG-3’) and foxq2_Rv1WT (5’-TGCAC TGGAC ATGTG CCTCT G-3’) and (ii) foxq2_Fw1mutA (5’-TGGCT AACAA GAGG GCAAT GTAGG-3’). PCR primers used for amplification are listed in table S3.

The mutant larvae and its siblings were dissected into anterior and posterior segments; the posterior parts were used for genotyping, while the anterior segments were soaked in RNAlater (Sigma-Aldrich) for RT-qPCR analysis or fixed with 4% paraformaldehyde (PFA) in Ca2+- and Mg2+-free Dulbecco’s phosphate-buffered saline (D-PBS) for cryosectioning.
set of PCR primer, which amplifies both rh2-1 and rh2-2 opsin genes (referred to here as rh2-1/2) or both lws1 and lws2 opsin genes (referred to here as lws1/2). Expression levels of all the transcript isoforms of thrb were measured in Fig. 1A, while, in the rest of the experiments, we measured expression levels of a transcript isoform of thrb, thrb2, which is essential for lws expression in mouse and zebrafish (13, 14).

In situ hybridization

In situ hybridization using ocular sections was carried out as described previously (17, 20). In short, zebrafish were anesthetized by chilling on ice and subjected to dissection the light phase of the light-dark cycle. The larval anterior segments or the adult eyes were fixed in 4% PFA in D-PBS overnight at 4°C. Before the fixation, the adult eyes were enucleated and poked with tweezers to make a tiny hole in the cornea. After sucrose infiltration and optimal cutting temperature compound embedding, the 10-μm frozen ocular sections were prepared with a cryostat. The cryosections were pretreated with proteinase K and hybridized with digoxigenin-labeled cRNA probes, and the hybridization signals were visualized by nitro blue tetrazolium–bromochloroindolyl phosphate staining. The images were acquired with an upright microscope (Axioplan2, Carl Zeiss). The cRNA probes were generated as described in our previous study (17, 20).

Immunofluorescent labeling of flat-mounted retina of adult zebrafish

Immunofluorescent labeling of flat-mounted retina was carried out as described (20) with some modifications. Briefly, retinas were isolated from dark-adapted adult zebrafish, and each of the retinas was processed to have four small radial cuts with equal spacing. Each retina was placed onto a piece of Parafilm sheet in a ~50-μl drop of fixative (4% PFA in D-PBS with 5% sucrose) and incubated at room temperature for 15 min. The fixed retina was then covered with another piece of Parafilm sheet over which a 1.5-g weight (1.5-ml tube containing water) was placed. After 30-min incubation at room temperature, the flattened retina was placed in 1 ml of fixative (4% PFA in D-PBS with 5% sucrose) and incubated at room temperature for 1 hour. The fixed retina was then washed with antibody diluent three times for 10 min each and treated with goat anti-mouse immunoglobulin G (IgG) antibody conjugated to Alexa Fluor 568 (diluted to 2 μg/ml; Molecular Probes, A-11004) in the antibody diluent for ~18 hours at room temperature. The retina was then washed with the antibody diluent three times for 10 min each and treated with goat anti-mouse immunoglobulin G (IgG) antibody conjugated to Alexa Fluor 568 (diluted to 10 μg/ml; Molecular Probes, A-11004) in the antibody diluent for ~18 hours at room temperature. This second antibody reaction was performed in the presence of 10 μM DRAQ5 (DR5050, BioStatus) for staining of the cell nucleus. The retina was then washed with the antibody diluent three times for 10 min each, mounted onto an agarose-coated slide glass with the photoreceptor layer facing down, and cover-slipped with VECTASHIELD Mounting Medium. Fluorescent images of the immuno-stained retina were captured with a confocal laser scanning microscope (TCS SP8, Leica).

Hybridization chain reaction

FISH (fluorescence in situ hybridization) was conducted by using HCR v3.0 technology (55) according to the manufacturer’s instruction (Molecular Instruments). Whole-mount larvae (3 days postfertilization (dpf) or 5 dpf) were fixed in 4% PFA in D-PBS with 0.1% Tween 20 overnight at 4°C. Flat-mounted retinas were prepared as described above. Probe sets were listed in data file S2. Two probe sets, opn1mw1UTR and opn1mw2UTR, were designed against 3′ untranslated region (3′UTR) to distinguish expressions of these two genes, while other probe sets were designed against coding sequences. Combinations of Alexa Fluor–conjugated hairpins and probe sets used for each experiment were described in table S5. Fluorescent images of the whole-mount larvae were imaged with a light sheet fluorescence microscopy (Z.1, Zeiss). For cone mosaic analysis in the larval stage, eyes were dissected from the animal after HCR. Whole larval eyes and flat-mounted retinas were incubated with 4′,6-diamidino-2-phenylindole (DAPI) (1 μg/ml) for nuclear staining. Whole larval eyes were embedded in 1% low melting point agarose and imaged with a confocal laser scanning microscope (FV3000, Olympus). Flat-mounted retinas were cover-slipped with PermaFluor Mountant (Thermo Fisher Scientific) and observed with a confocal laser scanning microscope (FV3000, Olympus).

The number of cone opsin positive cells in each fluorescent image was counted using Analyzed Particle function in ImageJ (Fiji) version 2.1.0. First, the central region of the retina was cropped from the image (87.63 μm by 87.63 μm). The trimmed images were converted to 8-bit black and white images (“Convert to Mask”). The binarized objects were filled (“Fill Holes”), and connected components were cut apart into separate ones (“Watershed”). After a threshold was selected, Analyze Particles settings were set to size 5 to 100 (μm2) and circularity 0.05 to 1.00. Six images from three individuals (two eyes each) for each genotype were processed for the quantification in Fig. 6E and fig. S7, while three images from three individuals were processed in Fig. 2E.

TUNEL staining

One head of an adult fish was subjected to cryosectioning for each genotype in a single experiment. Frozen ocular sections (10 μm thick) were prepared as described in the previous section for in situ hybridization. The cryosections were subjected to TUNEL staining by using the Click-IT TUNEL Alexa Fluor 594 Kit (Invitrogen) according to the manufacturer’s protocol. The stained sections were coverslipped with PermaFluor Mountant (Thermo Fisher Scientific) after incubation with DAPI (1 μg/ml) for nuclear staining, and observed with a confocal laser scanning microscope (FV3000, Olympus).

Immunohistochemistry

Immunohistochemistry with ocular sections was carried out as described previously (17, 20). Briefly, ocular cryosections were prepared as described in the previous section. The cryosections were pretreated with a blocking solution and then incubated with a primary antibody diluted in the blocking solution overnight at 4°C. After washed with PBS [10 mM Na-phosphate buffer, 140 mM NaCl, and 1 mM MgCl2 (pH 7.4)], the treated sections were immersed again with the blocking solution and then incubated for 4 hours at room temperature with a secondary antibody and with DAPI (3 μg/ml) for staining of the cell nuclei. The stained sections were coverslipped with VECTASHIELD Mounting Medium (Vector Laboratories) and imaged with a confocal laser scanning microscope (TCS SP8, Leica). The primary antibody used is mouse monoclonal antibody Zpr1 (diluted 1:400; Zebrafish International Resource Center, Eugene) against arrestin 3α. The secondary antibody used is goat anti-mouse IgG antibody conjugated with Alexa Fluor 568 (diluted to 2 μg/ml; Molecular Probes, A-11004).
Luciferase assay
For constructing the firefly luciferase reporter plasmids, the 1.56-kb sws2 upstream region with its 5′ UTR was amplified by PCR. The amplified fragment was ligated using the In-Fusion cloning kit (Takara) into the pGL4.13[luc2/SV40] (Promega, E6681) digested with Hind III and Bgl II. PCR primers used were as follows: IF-sws2-1.5kFwBgl (5′-GGAGG ATATC AGATC TAA CGG ATGGT TGCTG TTTGTG TGCTG-3′) and IF-sws2-RvHind (5′-CCCGA TTGCC AAGCT TTCCTG CATCGT AATTG GTGCC C-3′). The 0.53-kb sws2 reporter was constructed in a manner similar to the 1.56-kb sws2 reporter with following primers: IF-sws2_FwXho (5′-GCTCG CTAGC CTGCA GCAAC TGTA CTAAG TG-3′), and IF-sws2_RvHind. The 1.56-kb sws2 reporter was truncated by PCR to generate the 0.3- and 0.25-kb sws2 reporter by using the following PCR primers: pGL3-sws2_300_Fw (5′-TGGTT CACTG CGCAG ATGTA G-3′), 250sws2_Fw (5′-GAAAAC TTGGT GTGTA GCTGA TG-3′), and pGL4.13_Rv (5′-AGATC TGATA TCCTC GAGGC TAG-3′). For generating the pGL4 vector having no basal promoter, the SV40 promoter in the pGL4.13 vector was removed by a combination of enzyme digestion and self-ligation; the pGL4.13 vector was double digested with Hind III and Xho I, treated T4 DNA polymerase to make a blunt end, and self-ligated. The nucleotide mutations on the potential Foxq2-binding motif were introduced into the 0.25-kb SWS2 reporter by PCR. The promoter sequence in each of these resultant constructs was sequenced to confirm that no unintended mutation was incorporated into the promoter region.

To generate the expression plasmid of Foxq2, we first inserted a FLAG epitope tag into the pcAG vector (gifted from T. Matsuda). The resultant vector was named as pCAG-FLAG. We then amplified the cDNA fragments of foxq2 from retinal cdNAs of adult zebrafish and cloned them into the EcoRV-treated pCAG-FLAG vector. These plasmids were named as drFoxq2-pCAG-FLAG. For generating the VP64-Foxq2 expression plasmid, we first cloned the multiple repeats of the herpes simplex VP16 activation domain (synthesized DNA fragments) into the Xho I–treated pCAG-FLAG vector. The resultant plasmid, named as pCAG-FLAG-VP64N, was treated with the EcoRV and ligated with the PCR-amplified cDNA fragment. The nucleotide deletion in the DNA binding domain of Foxq2 was introduced by PCR. The DNA sequence for drFoxq2/pCAG-FLAG was constructed with a blunt end, and self-ligated. The nucleotide mutations on the potential Foxq2-binding motif were introduced into the 0.25-kb SWS2 reporter by PCR. The promoter sequence in each of these resultant constructs was sequenced to confirm that no unintended mutation was incorporated into the promoter region.

Immunoblot analysis
Immunoblot analysis was carried out as described previously (20). In short, proteins lysed in an SDS-PAGE sampling buffer were separated on a gel by SDS-PAGE, transferred to a Immobilon-P transfer membrane (Millipore), and probed with primary antibodies overnight at 4°C. The bound primary antibodies were detected by horseradish peroxidase–conjugated secondary antibodies in combination with an enhanced chemiluminescence detection system using Western Lightning Chemiluminescence Reagent (PerkinElmer Life Sciences) or ImmunoStar (Wako Pure Chemical Industries). Chemiluminescent images were acquired with ImageQuant LAS 4000 mini (GE Healthcare). The primary antibodies used were as follows: anti-FLAG antibody (diluted to 0.8 μg/ml; Sigma-Aldrich, F3165) and anti-H3K4me3 antibody (diluted 1:5000; Upstate, 07-473). The secondary antibodies used were as follows: horseradish peroxidase–conjugated anti-mouse IgG (diluted to 0.2 μg/ml; Kirkegaard & Perry Laboratories, 074-1816) and horseradish peroxidase–conjugated anti-rabbit IgG (diluted to 0.2 μg/ml; Kirkegaard & Perry Laboratories, 074-1516).

Generation of transgenic zebrafish
To construct a Foxq2 transgene plasmid, the FLAG-Foxq2 coding sequence was amplified by PCR from the plasmid, drFoxq2/pCAG-FLAG (described in the section of Luciferase assay). The amplified fragment was then cloned with an In-Fusion HD cloning kit into the pT2drCrx5.2kGP2ASix7 (20) digested by Xho I and Bam HI. The resultant plasmid, named pT2drCrx5.2kGP2Afoxq2, was used for the generation of transgenic zebrafish. The foxq2 transgenic zebrafish, ja78Tg, ja79Tg, and ja91Tg strains, were generated with the Tol2-based transgenesis system (56). The Tol2 transposase mRNA was transcribed from pCS-TP in vitro using the SP6 mMESSAGE mMACHINE Kit (Ambion) and purified with the RNEasy Mini kit (Qiagen). The purified Tol2 mRNA and the plasmid DNA were mixed and diluted to a final concentration of 25 ng/μl for each in 0.05% phenol red solution. About 1 nl of the DNA/RNA solution was injected into each of WT embryos at the one-cell stage. Fluorescence-positive embryos were isolated and raised to adulthood. The raised F0 founder fish were crossed with the WT fish, and subsequent F1 embryos were screened by the presence of fluorescence at four or 5 dpf. The transgenic lines were established from individual F0 fish. The foxq2-tg larval fish and its siblings were genotyped by ocular EGFP fluorescence just before the sampling and soaked in RNAi later for RT-qPCR analysis or fixed with 4% PFA in D-PBS for cryosectioning. The GFP-negative siblings were used as a control. To unambiguously identify the transgenic larvae according to ocular EGFP expression, synthesis of melanin pigment was inhibited by treating embryos with the egg water containing 0.003% 1-phenol-2-thiouracil from 24 hours to 5 days after fertilization.

Motif scanning
The sws2 promoter (fig. S6) was sequence-verified by traditional Sanger sequencing and used for the motif scanning. Fox and Crx binding profiles, each represented as a matrix consisting of nucleotide

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counts per position, i.e., position frequency matrix, were retrieved from the JASPAR CORE database (http://jaspar.genereg.net). These retrieved binding profiles (total of 11 profiles) were composed of one binding profile of Crx in mouse (57) and two binding profiles (FkhP and FkhS) for each of five Fox proteins in mouse (FoxA2, FoxL1, FoxK1, FoxJ1, and FoxJ3) (58). These frequency matrices were used to construct position-dependent letter-probability matrices that describe the probability of each possible letter at each position in the pattern with the simplest background model assuming that each letter appears equally frequently in the dataset. The sws2 promoter was scanned for individual matches to each of the Crx and Fox motifs with FIMO v5.1.1 (59). Biased distribution of individual letters in the promoter sequences was normalized by a zeroth-order model of Markov background probabilities constructed with the fasta-get-Markov tool in the MEME suite v5.1.1 (59) using nucleotide sequences between 1000-bp upstream and 1000-bp downstream of a transcription start site for all protein-coding genes in zebrafish (genome assembly GRCz11, Ensembl Release 98). All motif occurrences with $P < 1 \times 10^{-4}$ are indicated in fig. S6. The motif scoring results and matrix identities (IDs) of Fox and Crx in the database are included in data file S3.

BLAST searches and phylogenetic analysis
For identifying FOXQ2 genes in platypus and chicken, tBLASTn search (Ensembl web tools) was conducted against genome sequences of platypus (reference genome ID: mOrnAna1.p.v1) and chicken (reference genome ID: GRCg6a) using the amino acid sequence of forkhead domain of zebrafish Foxq2 (Ensembl protein ID: ENSDARP00000119225.2) as the query. We retrieved nucleotide sequences in intergenic regions showing the higher alignment score than any other regions encoding members of Fox families. The retrieved nucleotide sequences of FOXQ2 genes were mapped onto two distinct genomic regions but were adjacent in the region of the same chromosome, where any other gene is not annotated. We thus assumed that these mapped regions are two exons of FOXQ2 genes. Consistently, the forkhead domain of zebrafish Foxq2 is encoded in two exons. We then manually annotated exon-exon junctions of FOXQ2 genes according to the GT/AG mRNA processing rule. The annotated cDNA sequence of the forkhead domain of FOXQ2 gene was translated into a protein sequence and used for constructing a phylogenetic tree described in the next paragraph. The nucleotide sequences and annotations of FOXQ2 genes are summarized in data file S4.

For constructing a phylogenetic tree, FOXQ2 amino acid sequences in spotted gar, medaka, coelacanth, and sparrow hawk were retrieved from Ensembl database (Ensembl Release 101). FOXQ2 sequences in platypus and chicken identified by our blast searching were also used for the phylogenetic tree construction. Amino acid sequences of other members of Fox subfamilies were retrieved by performing tblastn searches (NCBI) against ReSeq RNA transcripts in purple sea urchin, zebrafish, chicken, platypus, and human using the amino acid sequence of forkhead domain of zebrafish Foxq2 as the query sequence with $E < 1 \times 10^{-20}$. Among the retrieved cDNA sequences, members of representative Fox subfamily (FOXA, FOXB, FOXC, FOXF, FOXJ), and FOXQ) were selected and translated into amino acid sequences. The resultant sequences of Fox proteins were aligned by multiple sequence alignment programs: G-INS-i program in MAFFT v7.471 under default settings (60). The aligned sequences were trimmed, remaining the sequences of the forkhead domain. Alignment gaps were manually inspected and deleted. The maximum likelihood tree was inferred by RAxML-NG v1.0.2 (61). The best tree was selected out of 40 alternative runs on 20 random- and 20 parsimony-based starting trees (–tree pars[20], rand[20] option). The amino acid replacement models of Le-Gascuel (LG) with gamma distribution (G4) were selected using the Akaike information criterion implemented in ModelTest-NG version x.y.z (62). The bootstrap values were obtained from sampling 500 times. The amino acid sequences used for the construction of phylogenetic tree are listed in data file S4. Accession numbers for genome assemblies and Fox genes are also provided in data file S4.

Statistical analysis
Sample sizes were determined on the basis of prior literature and best practices in the field, and no statistical methods were used to predetermine sample size. A two-tailed unpaired t test was used to determine the statistical significance between two datasets (Excel). Tukey-Kramer honestly significant difference test was used to determine the statistical significance among multiple datasets (the “stats” package in R, version 3.6.1).

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at https://science.sciencemag.org/content/8/1/eaib9784

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