Retinal Cell Biology

Retinal Cone Mosaic in sws1-Mutant Medaka (Oryzias latipes), A Teleost

Megumi Matsuo,1 Makoto Matsuyama,2 Tomoe Kobayashi,2 Shinji Kanda,3 Satoshi Ansai,4 Taichi Kawakami,5 Erika Hosokawa,5 Yutaka Daido,5 Takehiro G. Kusakabe,5 Kiyoshi Naruse,4 and Shoji Fukamachi1

1Department of Chemical and Biological Sciences, Japan Women’s University, Bunkyo-ku, Tokyo, Japan
2Division of Molecular Genetics, Shigei Medical Research Institute, 2117 Yamada, Minami-ku, Okayama, Japan
3Laboratory of Physiology, Atmosphere and Ocean Research Institute, The University of Tokyo, Kashiwa, Chiba, Japan
4Laboratory of Bioresources/NIBB Center of the Interuniversity Bio-Backup Project, National Institute for Basic Biology, Okazaki, Aichi, Japan
5Institute for Integrative Neurobiology and Department of Biology, Graduate School of Natural Science, Konan University, Kobe, Hyogo, Japan

Correspondence: Megumi Matsuo and Shoji Fukamachi, Department of Chemical and Biological Sciences, Japan Women’s University, Meijirdai 2-8-1, Bunkyo-ku, Tokyo 112-8681, Japan; mmatsuo@fc.jwu.ac.jp and fukamachi@fc.jwu.ac.jp.

Received: March 31, 2022
Accepted: October 2, 2022
Published: October 27, 2022

Citation: Matsuo M, Matsuyama M, Kobayashi T, et al. Retinal cone mosaic in sws1-mutant medaka (Oryzias latipes), a teleost. Invest Ophthalmol Vis Sci. 2022;63(11):21. https://doi.org/10.1167/iovs.63.11.21

Purpose. Ablation of short single cones (SSCs) expressing short-wavelength-sensitive opsin (SWS1) is well analyzed in the field of regenerative retinal cells. In contrast with ablation studies, the phenomena caused by the complete deletion of SWS1 are less well-understood. To assess the effects of SWS1 deficiency on retinal structure, we established and analyzed sws1-mutant medaka.

Methods. To visualize SWS1, a monoclonal anti-SWS1 antibody and transgenic reporter fish (Tg(sws1:mem-egfp)) were generated. We also developed a CRISPR/Cas-driven sws1-mutant line. Retinal structure of sws1 mutant was visualized using anti-SWS1, 1D4, and ZPR1 antibodies and coumarin derivatives and compared with wild type, Tg(sws1:mem-egfp), and another opsin (lus) mutant.

Results. Our rat monoclonal antibody specifically recognized medaka SWS1. Sus1 mutant retained regularly arranged cone mosaic as lus mutant and its SSCs had neither SWS1 nor long wavelength sensitive opsin. Depletion of sws1 did not affect the expression of long wavelength sensitive opsin, and vice versa. ZPR1 antibody recognized arrestin spread throughout double cones and long single cones in wild-type, transgenic, and sws1-mutant lines.

Conclusions. Comparative observation of sws1-mutant and wild-type retinas revealed that ZPR1 negativity is not a marker for SSCs with SWS1, but SSCs themselves. Loss of functional sws1 did not cause retinal degeneration, indicating that sws1 is not essential for cone mosaic development in medaka. Our two fish lines, one with visualized SWS1 and the other lacking functional SW1, offer an opportunity to study neural network synapsing with SSCs and to clarify the role of SWS1 in vision.

Keywords: SWS1 (short wavelength sensitive opsin), LWS (long wavelength sensitive opsin), arrestin, cone photoreceptor, retinal degeneration

Animals live in response to their environment. In vertebrates, the eyes play a major role in vision. In the retina of the eye, photoreceptor cells sensitive to various wavelengths convert light stimuli into cellular signals, which are then transmitted to downstream neurons.1 Teleosts possess rod cells and four subtypes of cone cells. Typically, rods and cones contain a single visual pigment,2–7 opsin protein bound to a chromophore, which together determines spectral sensitivity.8 There are four opsins in cones (short-wavelength-sensitive opsin [SWS1], SWS2, rhodopsin-like [RH], long-wavelength-sensitive opsin [LWS]), which are sensitive to UV, blue, green, and red light, respectively) (Fig. 1) and one in rods (RH1). Teleost cone cells form an ordered array with regular spacing, where red and green cones exist as physically fused double cones (DCs).9–12 The blue cone is long with a single outer segment (long single cone [LSC]). The UV cone is short with a single outer segment (short single cone [SSC]). DC, LSC, and SSC formed the crystalline arrays, which are generally categorized into two patterns: row mosaic and square mosaic.13–15 In row mosaics as of zebrasfish, double and single cone photoreceptors are arranged in parallel rows.16,17 In other teleost such as goldfish and medaka, retinal cone cells form a lattice arrangement of squares (Fig. 1)18–20 In rainbow trout, these two patterns of cones coexist in the retina.14,30

In both rod and cone cells, the light activates visual pigments, which subsequently induces a phototransduction cascade.21 The termination of light response requires inactivation of light-activated visual pigments by phosphorylation, and visual arrestin completely shuts off the activity of
light-activated visual pigments by binding to phosphorylated pigments. The functional null mutations in Arrestin result in a type of congenital stationary night blindness called Oguchi disease. The visual arrestin is a cytosolic protein with a molecular weight of 40 to 45 kDa, which is encoded by the genes SAG (rod arrestin, expressed in rods) and Arrestin3 (Arr3, cone arrestin, expressed in cones). Fish have two subtypes of Arr3, Arr3a and Arr3b. Labeling with subtype-specific antibodies (e.g., ZPR1 antibody for Arr3a) reveals subfunctionalized expression of Arr3a in DC and Arr3b in LSC and SSC in zebrafish.

We have used medaka (Oryzias latipes), a model animal with a long history of genetic research. Many established resources, laboratory strains (https://shigen.nig.ac.jp/medaka/top/top.jsp), and techniques used in biological, ecological, behavioral fields are available. Teleost visual pigments with LWS or SWS1 absorb the longest or shortest parts of wavelength, respectively. We established lws-mutant medaka and showed that decreased red light sensitivity affected behavioral response. SWS1-expressing SSCs work in prey capture, but in contrast with lws, the effects of the loss of sus1 have not been well-analyzed, except in mouse and trout. Although a lack of sus1 in mouse does not cause retinal degeneration, a recent study of CRISPR/Cas9 genome editing system (Fig. 2) as described previously. Primer sets to check the transcripts in

Thus, we investigated the effect of sus1 deficiency on retinal structure and arrestin expression using medaka. To this end, we established two medaka lines, a transgenic sus1-reporter and an sus1-deficient line, and produced a monoclonal antibody specifically recognizing SWS1.

**METHODS**

**Medaka Husbandry**

We used laboratory-raised, 3- to 12-month-old matured medaka (O. latipes). Fish were reared under a 14/10-hour light/dark cycle. All the treatments of animals in this research were carried out in accordance with the Japanese Act on Welfare and Management of Animals (Act No. 105 of October 1, 1973; the latest revisions Act No. 51 of June 2, 2017, effective June 1, 2018). All experimental protocols were approved by the Institutional Animal Care and Use Committees of Konan University and by the Animal Experiment Committees of Japan Women's University.

**Establishment of sus1-mutant Medaka**

Sus1-mutant medaka was generated with the CRISPR-Cas9 genome editing system (Fig. 2) as described previously.
The sws1-mutant medaka. (A) Genomic and translated peptide sequence of the medaka SWS1 locus (AB223058). Initiation and termination codons are underlined. Exons and introns are shown as black capital and gray small letters, respectively. Magenta indicates the target sequence for CRISPR/Cas9. The TMHMM predicted transmembrane regions are highlighted in yellow. The peptide sequence used as an antigen for antibody production (see Fig. 3) is highlighted in purple. (B) Electropherogram of the sws1 mutations. The target sequence is highlighted in magenta. These electropherograms are flipped (i.e., shown as a reverse complement) from the original ones. Purple horizontal lines in sws1+14 indicate tandem repeats of 15 nucleotides. (C) Deduced SWS1 peptide sequence of sws1−10, aligned with SWS1 of wild type. SWS1 of sws1−10 medaka was 88 amino acids long, which has an identical N-terminus to wild-type SWS1 and has the first of seven transmembrane regions of intact SWS1. (D) Offspring between the sws1 heterozygotes. 286 and 64 adults were genotyped for sws1−10 and sws1+14, respectively. Mendelian inheritance should result in a genotype ratio of wild-type:heterozygotes:homozygotes of 1:2:1, which was observed in both mutations (P = 0.482 for sws1−10 and 0.305 for sws1+14, χ² test). (E) RT-PCR. cDNA synthesized from mRNA in the eye (n = 3 each for wild type and sws1−10) was used as a template. Transcript of SWS1 was decreased in the mutant, whereas that of actin beta (Actb) was equivalent between the strains.

Generation of Anti-SWS1 Monoclonal Antibody

Opsin peptide sequences for medaka RH1 (BAD99136.1) were aligned using CLUSTAL-Omega at the European Molecular Biology Laboratory–European Bioinformatics Institute. Based on sequence comparison, the C-terminus of SWS1 peptide was chosen as immunogens for generating antibodies (Fig. 3A). We produced rat monoclonal anti-SWS1 antibody, as described previously. The supernatants of 5 of 39 hybridoma clone cultures were subjected to the immunohistochemical analysis described in this article.

Sexually matured d-rR strain medaka were kept in a dark room for 1 hour to make their melanin granules of pigment epithelium aggregate at the basal region of the cells. They were then deeply anesthetized with MS-222 (Sigma, St. Louis, MO) and perfused with 4% paraformaldehyde in 0.05 M PBS (pH 7.4) from the conus arteriosus. Retinas were dissected and post-fixed with the same fixative at least 1 hour at 4°C. They were immersed in 30% sucrose in PBS until well-soaked, embedded in 5% agarose (Sigma Type IX) solution containing 20% sucrose, and quickly frozen in cold n-hexane (−60°C). Cross-sections were prepared on slide glasses with a cryostat at 30 μm and dried at room temperature (RT). After penetrating with PBS containing 0.3% Triton X-100, the sections were incubated with one of the supernatants (10% supernatant for fluorescent detection or 2% for 3,3′-diaminobenzidine detection, 5% normal goat serum in PBS) overnight. They were reacted with biotinylated anti-rat IgG (Jackson ImmunoResearch,
West Grove, PA) for 1 hour, subsequently with ABC complex (VECTASTAIN ABC kit, vector lab, Burlingame, CA) for 1 hour. Finally, sections were visualized through the fluorescence of Streptavidin, Alexa Fluor 488 (Invitrogen, Waltham, MA) conjugate. In some samples, we performed 3,3′-diaminobenzidine detection followed by counterstaining with hematoxylin and eosin staining (HE). Cone-shaped SWS1 immunoreactive signals were observed in the inner/outer segment (IS/OS) layers. GCL, ganglion cell layer; IPL, inner plexiform layer; INL, inner nuclear layer; RPE/C, retinal pigment epithelium/cells.

Establishment of *sus1:mem-egfp* TG Line of Medaka

The 5′ flanking sequence of the medaka SWS1 gene obtained from the Ensembl genome database was used to design gene-specific primers. The *O. latipes* genomic DNA was extracted from one individual of the Hd-rR inbred strain as described previously.60 Genomic DNA fragments containing the upstream cis-regulatory region of SWS1 were amplified from the genomic DNA by PCR using a thermostable DNA polymerase (PrimeStar HS DNA polymerase, Takara BIO, Japan) and a pair of gene-specific oligonucleotide primers. 5′-TGACGTACGATCTCGTTCTGTTCTGCTCTG-3′ and 5′-ACGGATCCGATCTCGTTCTGTTCTGCTCTG-3′.62 The pBluescript-mem-EGFP vector was made by inserting the oligonucleotides corresponding with the membrane-anchoring signal of neuremodulin (5′-atgctgtgctgtatga-3′) into the 5′ end of the EGFP-coding region of pBluescript-EGFP, generating an open reading frame encoding a fusion protein of the N-terminal 20 amino acids of neuremodulin and EGFP that can label plasma and intracellular membranes with fluorescence.64 The amplified 1.2-kb upstream region of SWS1 was inserted into the *SalI/BamHI* sites of pBluescript-mem-EGFP. The resultant plasmid DNA was introduced into medaka embryos by microinjection as described previously.65 The injected embryos were screened for fluorescence in retinal photoreceptor cells under a fluorescent dissection microscope (M165 FC; Leica Microsystems, Wetzlar, Germany) and then reared to adult fish. The transgenic strain *Tg(sus1:mem-egfp)* was established by successive crosses and selection of the fish with green fluorescent protein (GFP) fluorescence in retinal photoreceptor cells. In immunohistological analyses, for double labeling of the *Tg(sus1:mem-egfp)* retina with GFP and anti-SWS1 or ZPR1 (Abcam, Cambridge, UK), GFP was visualized by immunostaining with rabbit anti-GFP polyclonal antibody (Invitrogen) and Alexa Fluor 488-conjugated anti-rabbit IgG secondary antibody, diluted 1000-fold. Anti-SWS1 and ZPR1 signals were detected with Alexa Fluor 594-conjugated anti-rabbit IgG secondary antibody, diluted 1000-fold. Anti-SWS1 and ZPR1 signals were detected with Alexa Fluor 594-conjugated anti-rabbit IgG secondary antibody, diluted 1000-fold. Anti-SWS1 and ZPR1 signals were detected with Alexa Fluor 594-conjugated anti-rabbit IgG secondary antibody, diluted 1000-fold. Anti-SWS1 and ZPR1 signals were detected with Alexa Fluor 594-conjugated anti-rabbit IgG secondary antibody, diluted 1000-fold.

Histology of Whole-mount Retina

The whole retina was isolated from the eyecup of dark-adapted medaka according to the method of Salbreux et al.,66 with some modification. The retina was flushed out from the enucleated eye and rinsed in ice-chilled PBS. After the isolation and fixation in 4% paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) with 5% sucrose for 15 minutes twice and 30 minutes once, the retina was transferred onto nonfluorescent adhesive glass slides (MAS-coated glass slides, Matsunami Glass Ind., Osaka, Japan). The fixed retina was rinsed three times with 0.1M phosphate buffer containing 5% sucrose every 20 minutes and then once with PBST (PBS with 1% Tween, 1% Triton X-100, and 1% DMSO) for Figure 3. Establishment of an anti-SWS1 monoclonal antibody. (A) The peptide sequence of medaka SWS1 used for immunization to rats (purple) is aligned with the sequence of other opsins of medaka. The color of amino acid residues means their physicochemical properties. An asterisk (*) indicates a position that has a single, fully conserved residue. A period (.) indicates conservation between groups of weakly similar properties. All the settings are the defaults for Clustal Omega. (B) Establishment of an anti-SWS1 monoclonal antibody. (A) The peptide sequence of medaka SWS1 used for immunization to rats (purple) is aligned with the sequence of other opsins of medaka. The color of amino acid residues means their physicochemical properties. An asterisk (*) indicates a position that has a single, fully conserved residue. A period (.) indicates conservation between groups of weakly similar properties. All the settings are the defaults for Clustal Omega.

Establishment of an anti-SWS1 monoclonal antibody. (A) The peptide sequence of medaka SWS1 used for immunization to rats (purple) is aligned with the sequence of other opsins of medaka. The color of amino acid residues means their physicochemical properties. An asterisk (*) indicates a position that has a single, fully conserved residue. A period (.) indicates conservation between groups of weakly similar properties. All the settings are the defaults for Clustal Omega.
15 minutes. After rinsing, the retina was blocked in 10% normal rabbit serum at RT for 1 hour, treated with anti-SWS1 antibody diluted 10 times by PBST, covered with dice-size-cut parafilm in a moist chamber, and incubated at RT for more than 14 hours. The specimen was washed twice with PBST for 10 minutes, then once with PBS for 5 minutes. These washing steps were conducted routinely between each step. After 2 hours of incubation with biotinylated anti-rat IgG, followed by ABC reagents for 30 minutes, Alexa Fluor 555-conjugated streptavidin (1/1000) was reacted for 30 minutes at RT. Finally, the retina was covered with ProLong Gold (Invitrogen).

For immunocytochemistry with ZPR1 or 1D4 (ab54177, Abcam, UK), each antibody in PBS (1/1000) with 5% normal rabbit serum was reacted at RT for more than 14 hours. The retina was incubated in PBS with rabbit anti-mouse IgG conjugated with Alexa Fluor 633 (1/500, Invitrogen) for 1 hour at RT. We injected 3-(2-Benzothiazolyl)-7-(diethylamino)-coumarin (BTDEC, also known as coumarin 6; Tokyo Chemical Ind., Tokyo, Japan) into a secondary antibody solution (1 μg/mL) to label the retinal cell. Positively stained structures were pseudocolored in orange. For triple labeling using anti-SWS1, ZPR1 or 1D4, and BTDEC, a binding step of anti-SWS1 followed the incubation with ZPR1 or 1D4. Positive signals of anti-SWS1, 1D4, and ZPR1 were pseudocolored in violet, white, and cyan, respectively.

**Imaging of Whole-mount Retina**

The retinal images were recorded sequentially using Olympus FV1200 Laser Scanning Confocal Microscope (Olympus, Tokyo, Japan). Image stacks with 0.1 to 2.0 μm step depths were processed with FIJI software. Starting from the outer plexiform layer, z-depth increased toward the RPE.

**RESULTS**

**Establishment of sus1-mutant Line and Anti-SWS1 Monoclonal Antibody**

We previously established colorblind medaka by knocking out the *cone-opsin* genes using CRISPR/Cas9 technology. Using the same protocol but different guide RNA, we introduced frameshift mutations in the *SWS1* gene. The target sequence existed in the first exon coding the first transmembrane domain, and a codon for the second methionine was found in the second exon coding the fourth transmembrane domain (Fig. 2A). The frameshift mutations were a ten-base deletion (*sus1*−10) or a 14-base insertion (*sus1*+14) (Fig. 2B). The *sus1*+14 mutation had 15-base tandem repeats, which might reflect an error in the microhomology-mediated end joining. Opsins fold into a seven-transmembrane structure, typical of G protein-coupled receptors, but SWS1 of *sus1*−10 had only the first transmembrane region and was no longer a G protein-coupled receptor. Therefore, no functional protein could be translated from the mutated allele (Fig. 2C).

Heterozygotes or homozygotes of the *sus1* mutant were indistinguishable from wild type by appearance and seemed to be fully viable under our breeding condition (Fig. 2C) ($P > 0.05$, $\chi^2$ test). However, fertilized eggs could hardly be obtained from the *sus1*+14 homozygotes for an unknown reason; inbreeding depression might occur because of the limited number of offspring, and we gave up maintaining *sus1*+14 fish. Frozen sperms of the *sus1*−10 and *sus1*+14 mutants are available at NBRP medaka as MT1326 and MT1327, respectively. Transcripts of *sus1* were greatly reduced in the eyes of *sus1*−10 fish, likely reflecting nonsense-mediated mRNA decay (Fig. 2D). Our rat anti-SWS1 monoclonal antibody showed characteristic cone-shaped outer segments of cone cells in wild-type retina, but not in *sus1*−10 (Fig. 3).

**Retinal Cone Mosaic of Tg(sus1:mem-egfp) Medaka**

We visualized cone cells expressing *sus1* by generating a transgenic line, *Tg(sus1:mem-egfp)*. The 1.2-kb upstream region of medaka *sus1* containing cis-regulatory sequences was sufficient to drive reporter expression in cone cells (Fig. 4B, C). When a cone expressing *sus1* and having one protrusion was visualized with an anti-GFP antibody, its inner segment appeared as a green round and its outer segment as a green dot (Fig. 4C). Double immunolabeling of flat-mounted *Tg(sus1:mem-egfp)* retina with anti-GFP and ZPR1 antibodies revealed cone mosaics where a single cone subtype had GFP, and the other cone subtypes were labeled by ZPR1 (Fig. 4B). GFP-positive and ZPR1-positive cones together constituted square mosaics. When *Tg(sus1:mem-egfp)* retina was reacted with anti-SWS1 and anti-GFP antibodies, two signals colocalized in the outer segment of single cones but did not overlap in the inner parts of cells (Fig. 4C–E, Supplementary Video S1), suggesting that anti-SWS1 antibody specifically recognized the outer segments of single cones expressing *sus1*.

**Square Cone Mosaic Visualized Using Anti-SWS1, ZPR1 Antibodies, and Coumarin in Wild-type Medaka**

Next, we validated an arrangement of retinal cones in wild-type medaka, using coumarin derivatives and antibodies. Because coumarin stains retinal cells in zebrafish, we used BTDEC for immunohistochemistry of the whole mounted retina of medaka. BTDEC visualized cone cells spread over the retina (Fig. 5), where fluorescence was strong in the outer segment and somewhat weak in the inner segment of cones (Fig. 6). ZPR1 visualized cone photoreceptors, from the tip of the outer segment to the inner segment, axon, and synaptic terminal (Fig. 1). The retina with ZPR1 and anti-SWS1 antibodies showed that ZPR1-negative cones had SWS1 monoclonal antibody showed characteristic cone cells expressing anti-SWS1 antibodies, two signals colocalized in the outer segment of single cones but did not overlap in the inner parts of cells (Fig. 4C–E, Supplementary Video S1), suggesting that anti-SWS1 antibody specifically recognized the outer segments of single cones expressing *sus1*.

**Retained Regularity of Cone Arrangement in sus1−10 Medaka’s Retina**

To assess the effects of SWS1-depletion on retinal structure, we analyzed *sus1*−10 medaka using the same strategies
FIGURE 4. Retinal cones of Tg(sws1:mem-egfp) medaka. (A) Schematic diagram showing the reporter construct of sws1:mem-egfp. Numbers in parentheses indicate the nucleotide positions relative to the transcription start site of sws1. An anti-GFP antibody visualized the retinal cones of Tg(sws1:mem-egfp) medaka. (B–D) Cone mosaic arrangement in Tg(sws1:mem-egfp) medaka’s retina. (B) Cells that expressed sws1 were visualized through the anti-GFP antibody. ZPR1 detected cells without anti-GFP labeling, LSCs and both members of DCs. An anti-GFP-positive cell was surrounded by ZPR1-positive cone cells, and both together made up a square mosaic. White arrows point to single cones, and yellow arrows point to DCs. (C) Anti-GFP signals were observed as green rounds and dots; that is, the inner segments and outer segments, respectively. (D) Anti-SWS1 signals were observed as violet dots; that is, outer segments. (E) Merged image of (B) and (C). Green dots of anti-GFP and violet dots of anti-SWS1 colocalized, showing that the outer segment with SWS1 was detected by a monoclonal antibody against SWS1. (B–D) White arrowheads point to anti-GFP-labeled green rounds. Asterisks point to cells in which both anti-GFP and anti-SWS1 signals overlapped.

as transgenic and wild-type medaka. The sws−10 medaka lost SWS1-signal in the retina, where BTDEC-stained cones formed regular arrangement (Fig. 7) just as seen in wild-type medaka (Fig. 5). At depths where a single outer segment of a ZPR1-negative cell appeared, inner segments of other cone subtypes were observed as orange rounds (Fig. 7).

A z-depth stack model visualized retinal cells of sws1−10 medaka; BTDEC stained both rods and cones, whereas ZPR1 stained cones (Fig. 8A). The stacks of z = 0 to 15.6 μm with BTDEC and ZPR1 showed the outer and inner segments of cones. The stacks of z = 0 to 8.8 μm exhibited the inner segments, forming square mosaics (Fig. 8A). Higher magnified images represented that the outer segment of a ZPR1-negative cell was surrounded by the inner segments of four DC cells and four single cone cells (Fig. 8B). An orthogonal view of the z-stacks of BTDEC and ZPR1 showed the entire cone (Fig. 8C). All cones were visualized with BTDEC, regardless of ZPR1 signal. ZPR1-negative cones were shorter than the surrounding cones (Fig. 8C). Comprehensive considering Figures 7 and 8, in sws1−10, ZPR1-negative cells were SSCs. Overall, sws1−10 medaka retained...
FIGURE 5. Retinal cones of wild-type medaka. Photoreceptor cells in medaka’s retina were visualized with ZPR1 (cyan) and BTDEC (orange) in the upper panel, and with anti-SWS1 (violet), ZPR1 antibodies, and BTDEC in the lower panel. BTDEC showed the regular arrangement of cone cells where the ZPR1 signal was absent in one cone subtype. Violet anti-SWS1 signal existed in the ZPR1-vacant space, showing that the cone cell expressing SWS1 was ZPR1-negative.

FIGURE 6. Volume-rendered z-stacksof the retinal photoreceptor cells of wild-type medaka. (A) Photoreceptor cells visualized by BTDEC and anti-SWS1 antibody. Shown are fluorescent images (upper panel) and z-depth stacks color-coded for depth (lower panel). BTDEC visualized rod photoreceptor cells and regularly arranged cone cells. The Inner segments of cones were observed at deeper positions than the outer segments. (B) A yellow boxed retinal area in (A) shown at higher magnification. A single protruding outer segment with SWS1 was blue in z-stacks. The cone with SWS1 was surrounded by other cones whose outer segments were magenta to red. Thus, cone cells expressing SWS1 were shorter than other cone subtypes.

regular cone mosaic arrangements just as wild type. Even though *sus1<sup>−10</sup>* did not express SWS1, SSC was not lost. Furthermore, in *sus1<sup>−10</sup>* medaka’s retina, ZPR1 labeled both DC and LSC, but not SSC.

Next, we analyzed previously established another *opsin*-gene mutant, *lus<sup>1-Δ11Δ5b</sup>* line, to compare the effect of opsin depletion on retinal development. Anti-SWS1 antibody visualized cells with regular spacing, and BTDEC merged
FIGURE 7. Retinal cones of *sws1*−10 medaka. A single dot-shaped outer segment of a ZPR1-negative cell was surrounded by round-shaped inner segments of other cones.

mosaic arrangement of cones in *lws*+2a+5b retina (Fig. 9). The 1D4 antibody binds to bovine rhodopsin,71 but in zebrafish, it labels LWS-expressing cells,34 and is used as a red cone marker.72 In wild-type medaka, 1D4 labeled the outer segment of one member of DC (Fig. 10A). The 1D4 signals disappeared in *lws*+2a+5b medaka (Fig. 9), revealing that it bound to LWS in DCs of medaka. Anti-SWS1 antibody labeled single cone of *lws*+2a+5b but not DCs (Fig. 9), and 1D4 conversely labeled one of the DCs in *sws1*−10 but not SSC (Fig. 10B). *Sws1* depletion had a negligible effect on *lws* expression and vice versa. In summary, neither retinal cone mosaic nor ZPR1- or 1D4-positive cells changed in *sws1*−10.

**DISCUSSION**

ZPR1 antibody binds to zebrafish Arr3a in DCs,33–36 which is orthologous to medaka Arr3a in DC and LSC.31 Given that DCs and LSCs were ZPR1 positive in this study, ZPR1 detected Arr3a in medaka. SSCs did not possess Arr3a, whether they expressed SWS1 or not. Thus, Arr3a negativity is a marker for SSCs, not for cones expressing SWS1. In zebrafish retina, the expression of arrestin is partitioned...
Retinal Cone Mosaic in sws1-Mutant Medaka

FIGURE 9. Retinal cones of lws1+2a+5b medaka. (A) An isolated retina observed under low magnification. Positively labeled cells with anti-SWS1 antibody spread over the retina, but no signal was obtained with 1D4 antibody. DT, light field image by differential interference observation; ON, optic nerve. (B) Two boxed areas in (A) observed under high magnification. Violet cells with anti-SWS1 antibody spread with regular spacing. (C) A retinal mosaic arrangement of cones. Only single cones were positively labeled with anti-SWS1 antibody. (D) Two images at identical x–y coordinates, but different depths. The outer segments of single cone cells had anti-SWS1 signal (upper panel), but DCs were not labeled by anti-SWS1 antibody (lower panel; 1.4 μm closer to the RPE than the upper panel). The yellow arrows indicate DCs, and white arrows indicate single cone cells.

among cone subtypes; DCs express Arr3a, whereas LSCs and SSCs express Arr3b.29 Contrasting zebrafish LSC, which possess Arr3b, medaka LSC had Arr3a. Whereas zebrafish and medaka have regularly arranged cones, LSCs of these two teleost species differ in the position of their mosaics, which may be related to the fact that the LSCs of these two species express different arrestin subtypes.

Mutations in blue, green, and red opsin and rhodopsin genes cause eye disorders and affect visual ability in humans.73–75 So far, animal models have been analyzed, indicating that the effects of mutations in opsin genes are not canonical and are diverse.76 For instance, in mice with a targeted disruption of rhodopsin, not only rods but also cones degenerate, and the outer segments of cones almost completely disappear.77 In zebrafish, meanwhile, mutations in rhodopsin gene cause rod degeneration, but cones are unaffected.78 In mice having mutations in M-opsin, M-opsin–dominant cones remain viable for at least 15 months, albeit with shortened or no outer segments, whereas S-opsin–dominant cones are normal.79,80 Given this finding, it would be interesting to know what opsins are expressed in the SSC of sus110 and DC of lws1+2a+5b medaka, both of which had neither SWS1 nor LWS. Besides histological observation, we assessed the behavioral phenotypes of sus1-mutant (SF, in preparation). Previously, we showed that lws- or sus2-mutant had reduced body color preferences.49 According to optomotor response, lws-mutant decreased red-light sensitivity46,48; however, blue light sensitivity did not change in sus2-mutant.55 We conducted mate choice and optomotor response assay of sus1-mutant, but their body color preference and UV sensitivity were equivalent to wild type. Furthermore, sus1/sus2–double-mutant had the same UV sensitivity as wild type. Considering these behavioral results, a new assay needs to be developed to assess the potential defect caused by the loss of SWS1 in medaka.

As mentioned elsewhere in this article, zebrafish is another well-studied fish model, but an sws1-mutant line has not been reported. Instead, ablation studies are available. Ablating zebrafish SSCs (expressing SWS1) stimulates a regenerative response.81–83 Zebrafish H3 horizontal cells exclusively connect to SSCs. When SSCs were ablated, H3 horizontal cells prioritize wiring with SSCs. However, when regeneration of SSCs is delayed or absent, H3 cells increase connections with LSC and DCs.82,84 It is unclear whether the morphology of SSCs or SWS1 of SSCs is important for horizontal cells to prefer SSCs. Elucidating the neural network postsynaptic to SSC of the sws1−10 retina will give us a clue to this question and suggest a role of SSCs without SWS1 in vision. So far, sws1-deficient animal models have been established in mouse and trout. The lack of sus1 does not cause retinal degeneration in mouse,52,53 but causes serious
FIGURE 10. Retinal cones of wild-type and sus1<sup>−10</sup> medaka. (A) Photoreceptor cells visualized by anti-SWS1, 1D4 antibodies and BTDEC in wild-type medaka. When a retina was rinsed well with PBS, few cones remained on the retina and each cone cell was easily identified. Anti-SWS1 antibody bound to a single cone cell, and 1D4 antibody did to one of DCs. (B) Z-stack images of retinal cones of sus1<sup>−10</sup> medaka. Shown are the z-stacks of z = 0 to 12.6 μm in the upper panel, and the z-stacks of z = 0 to 14.2 μm in the lower panel, accompanied by cartoons of cones on the right side. One member of DCs was labeled with 1D4 antibody, and the single cones were not.

retinal developmental defects in rainbow trout. Superscripted<br>Deficiency did not lead to retinal degeneration in medaka. The reason why the loss of sus1 did not induce serious effects as trout is unclear, but the sequence of opsins genesis may explain it. Lateral induction mechanisms have been anticipated as causal in creating fish cone mosaics in cyprinid fishes (goldfish and zebrafish), where the order of cone opsin appearance is LWS, followed by RH2, SWS1, and SWS2. In contrast, in situ hybridization results indicate that in salmonids, SWS1 appears first, followed by LWS, RH2, and SWS2. As per the lateral induction mechanism, defects of cone subtype induced early in retinal development would have a significant impact on the developing retina, whereas defects of cone subtype induced later would have little effect. Because SWS1 is the first opsin induced in trout, this mechanism sounds acceptable. If so, in medaka, SWS1 must be induced later in the developing retina, because SWS1 depletion did not affect retinal cone mosaics. Another interpretation of this study is that SWS1 is only a marker for wild-type SSCs in medaka. Because lws<sup>−24</sup> also retained its mosaic structure, opsins may not have a significant effect on retinal development in medaka. However, to determine whether these two hypotheses are plausible, further experiments are needed, such as elucidating the order of opsin genesis in medaka and analyzing the retinal structure of other opsin gene mutants.

CONCLUSIONS

Overall, we provided a monoclonal antibody specifically binding to medaka SWS1, sus1-reporter TG (Tg(sus1:mem-egfp)) line, and sus1-mutant line of medaka. Arr3a negativity was not a marker of cones with SWS1, but SSCs. Depletion of sus1 or lws did not cause retinal degeneration and did not affect each other’s expression. Our two fish models visualizing sus1 or lacking functional sus1 offer an opportunity to analyze the role of SSC in vision and neural networks postsynaptic to SSC.

Acknowledgments

The authors thank Kae Akita at JWU for helpful advice on the manipulation of the laser scanning confocal microscope, Maki Takahashi at JWU for RT-PCR of SWS1, and undergraduate students at JWU for rearing sus1-mutant and wild-type medaka. We also thank Shoji Kawamura at the University of Tokyo for his advice on epitope selection in producing anti-SWS1 antibodies, Ikuyo Hara at NIBB for establishing sus1-mutant and wild-type medaka, and Yuko Nishiwaki at OIST for critical reading of the article.

Supported by a Grant-in-Aid for Scientific Research (C) (17K07506) from the Japan Society for the Promotion of Science (JSPS), a grant for Joint Research (01111904) by the National Institutes of Natural Sciences (NINS), and research funds from JWU to S.F., Grants-in-Aid for Scientific Research from JSPS to KN (#16H04819), to T.G.K. (19H03213, 21K19280), and research grants from the Takeda Science Foundation and the Hirao Taro Foundation of KONAN GAKUEN for Academic Research to T.G.K.

Contributions: Mak.M and T.Ko. developed the rat anti-medaka SWS1 monoclonal antibody. S.K. screened and chose useful monoclonal antibodies specifically binding to the retinal photoreceptor layer of medaka. T.G.K. directed all the experiments of Tg(sus1:mem-egfp) medaka, and E.H. and Y.D. developed, and T.Ka. analyzed Tg(sus1:mem-egfp) medaka. K.N. and S.A. established sus1-mutant medaka. S.F. supervised the experiments using sus1-mutant and wild-type medaka. Meg.M analyzed sus1<sup>−10</sup>, lws<sup>−24</sup> and wild-type medaka and wrote...
References

1. Brzezinski JA, Reh TA. Photoreceptor cell fate specification in vertebrates. Development. 2015;142(19):3263–3273, doi:10.1242/dev.127043.

2. Shichida Y, Matsuyama T. Evolution of opsins and phototransduction. Philos Trans R Soc B Biol Sci. 2009;364(1531):2881–2895, doi:10.1098/rstb.2009.0051.

3. Imamoto Y, Shichida Y. Cone visual pigments. Biochim Biophys Acta Bioenerg. 2014;1837(5):664–673, doi:10.1016/j.bbabio.2013.08.009.

4. Owens GL, Rennison DJ, Allison W, Taylor JS. In the four-eyed fish (Anableps anableps), the regions of the retina exposed to aquatic and aerial light do not express the same set of opsin genes. Biol Lett. 2012;8(1):86–89, doi:10.1098/rsbl.2011.0582.

5. Takechi M, Kawamura S. Temporal and spatial changes in the expression pattern of multiple red and green subtype opsin genes during zebrafish development. J Exp Biol. 2005;208(7):1357–1345, doi:10.1242/jeb.01532.

6. Rennison D, Owens G, Allison W, Taylor J. Intra-retinal variation of opsin gene expression in the guppy (Poecilia reticulata). J Exp Biol. 2011;214(19):3248–3254, doi:10.1242/jeb.057836.

7. Stenkamp DL, Hisatomi O, Barthel LK, Tokunaga F, Raymond PA. Temporal expression of rod and cone opsins in embryonic goldfish retina predicts the spatial organization of the cone mosaic. Invest Ophthalmol Vis Sci. 1996;37(2):363–376.

8. Stenkamp DL, Barthel LK, Raymond PA. Spatiotemporal coordination of rod and cone photoreceptor differentiation in goldfish retina. J Comp Neurol. 1997;382:272–284.

9. Viets K, Eldred K, Johnston RJ. Mechanisms of photoreceptor patterning in vertebrates and invertebrates. Trends Genet. 2016;32(10):638–659, doi:10.1016/j.tig.2016.07.004.

10. Hunt DE, Rawlinson NJF, Thomas GA, Cobcroft JM. Investigating photoreceptor densities, potential visual acuity, and cone mosaics of shallow water, temperate fish species. Vision Res. 2015;111(1):13–21, doi:10.1016/j.visres.2015.03.017.

11. Cameron DA, Easter SS. Cone photoreceptor regeneration in adult fish retina: phenotypic determination and mosaic pattern formation. J Neurosci. 1995;15(3):2255–2271.

12. Carl HE, George Daniel S. The mosaic of single and twin cones in the retina of fishes. Am Nat. 1990;54(398):109–118, http://www.journals.uchicago.edu/t-an.

13. Flamarique IN, Cheng CL, Bergstrom C, Reimchen TE. Pronounced heritable variation and limited phenotypic plasticity in visual pigments and opsin expression of threespine stickleback photoreceptors. J Exp Biol. 2013;216(4):656–667, doi:10.1242/jeb.078840.

14. Cheng CL, Flamarique IN. Chromatic organization of cone photoreceptors in the retina of rainbow trout: single cones reversibly switch from UV (SWS1) to blue (SWS2) light sensitive opsin during natural development. J Exp Biol. 2007;210(23):4123–4135, doi:10.1242/jeb.009217.

15. Dalton BE, Loew ER, Cronin TW, Carleton KL. Spectral tuning by opsin coexpression in retinal regions that view different parts of the visual field. Proc R Soc B Biol Sci. 2014;281(1797):1–9, doi:10.1098/rspb.2014.1980.

16. Allison WT, Barthel LK, Skebo KM, Takechi M, Kawamura S, Raymond PA. Ontogeny of cone photoreceptor mosaics in zebrafish. J Comp Neurol. 2010;519(20):41825–4195, doi:10.1002/cne.22447.

17. Raymond PA, Barthel LK. A moving wave pattern of the cone photoreceptor mosaic array in the zebrafish retina. Int J Dev Biol. 2004;48(8–9):935–945, doi:10.1387/ijdb.041873pr.

18. Raymond PA, Barthel LK, Roussifere ME, Sullivan SA, Knight J. Expression of rod and cone visual pigments in goldfish and zebrafish: a rhodopsin-like gene is expressed in cones. Neuron. 1995;10(6):1161–1174, doi:10.1016/0896-6273(95)90064-X.

19. Nishiwaki Y, Oishi T, Tokunaga F, Morita T. Three-dimensional reconstitution of cone arrangement on the spherical surface of the retina in the medaka eyes. Zoolog Sci. 1997;14(5):795–801, doi:10.2108/zsj.14.795.

20. Hisatomi O, Satoh T, Tokunaga F. The primary structure and distribution of killifish visual pigments. Vision Res. 1997;37(22):3089–3096, doi:10.1016/S0042-6990(97)00115-6.

21. Kawamura S, Tachibanaki S. Rod and cone photoreceptors: molecular basis of the difference in their physiology. Comp Biochem Physiol A Mol Integr Physiol. 2008;150(4):369–377, doi:10.1016/j.cbpa.2008.04.600.

22. Chen CK, Burns ME, Spencer M, et al. Abnormal photoreceptors and light-induced apoptosis in rods lacking rhodopsin kinase. Proc Natl Acad Sci USA. 1999;96(7):3718–3722, doi:10.1073/pnas.96.7.3718.

23. Kühn H, Hall SW, Wilden U. Light-induced binding of 48kDa protein to photoreceptor membranes is highly enhanced by phosphorylation of rhodopsin. FEBS Lett. 1984;176(2):475–478.

24. Wilden U, Hall SW, Kühn H. Phosphodiesterase activation by photoexcited rhodopsin is quenched when rhodopsin is phosphorylated and binds the intrinsic 48-kDa protein of rod outer segments. Proc Natl Acad Sci USA. 1986;83(5):1174–1178, doi:10.1073/pnas.83.5.1174.

25. Xu J, Dodd RL, Makino CL, Simon MI, Baylor DA, Chen J. Prolonged photoreponses in transgenic mouse rods lacking arrestin. Nature. 1997;389(6650):505–509, doi:10.1038/39068.

26. Oguchi C. Über eine Abart von Hemeralopie. Acta Soc Ophthalmol Jpn. 1907;11:123–134.

27. Fuchs S, Nakazawa M, Maw M, Tamai M, Oguchi Y, Gal A. A homozygous 1–base pair deletion in the arrestin gene is a frequent cause of Oguchi disease in Japanese. Nat Genet. 1995;10(3):360–362, doi:10.1038/ng0795-360.

28. Gurevich VV, Hanson SM, Song X, Vishnevitskiy SA, Gurevich E V. The functional cycle of visual arrestins in photoreceptor cells. Prog Retin Eye Res. 2011;30(6):405–430, doi:10.1016/j.preteyeres.2011.07.002.

29. Renninger SL, Gesemann M, Neuhauss SCF. Cone arrestin confers cone vision of high temporal resolution in zebrafish larvae. Eur J Neurosci. 2011;33(4), doi:10.1111/j.1460-9586.2010.07574.x.

30. Imanishi Y, Hisatomi O, Tokunaga F. Two types of arrestins expressed in medaka rod photoreceptors. FEBS Lett. 1999;461(2–3):31–36, doi:10.1016/S0014-5793(9991483-0).

31. Ogawa Y, Corbo JC. Partitioning of gene expression among zebrafish photoreceptor subtypes. Sci Reports. 2021;11:17340, doi:10.1038/s41598-021-96837-z.

32. Hisatomi O, Imanishi Y, Satoh T, Tokunaga F. Arrestins expressed in killifish photoreceptor cells. FEBS Lett. 1997;411(1):12–18, http://www.ncbi.nlm.nih.gov/pubmed/9247134.

33. Larson KD, Bremiller R. Early onset of phenotype and cell patterning in the embryonic zebrafish retina. Development. 1990;109:567–576.
34. Yin J, Brocher J, Linder B, et al. The ID4 antibody labels outer segments of long double cone but not rod photoreceptors in zebrafish. Invest Ophthalmol Vis Sci. 2012;53(10):5480–5485, doi:10.1167/iovs.12-9511.

35. Raymond PA, Colvin SM, Jabeen Z, Nagashima M, Barthel LK. Patternning the cone mosaic array in zebrafish retina requires specification of ultraviolet-sensitive cones. PLoS One. 2014;9(1):85325, doi:10.1371/journal.pone.0085325.

36. Tappeiner C, Balmer J, Iglicki M, Schuerch K, Jazwinska A. Characteristics of rod regeneration in a novel zebrafish retinal degeneration model using N-methyl-N-nitrosourea (MNU). PLoS One. 2013;8(8):71064, doi:10.1371/journal.pone.0071064.

37. Kirchmaier S, Naruse K, Wittbrodt J, Loosli F. The genomic and genetic toolbox of the teleost medaka (Oryzias latipes). Genetics. 2015;199(4):905–918, doi:10.1534/genetics.114.173849.

38. Wittbrodt J, Shima A, Schartl M. Medaka - a model organism from the Far East. Nat Rev Genet. 2002;3(1):53–64, doi:10.1038/nrg704.

39. Kasahara M, Naruse K, Sasaki S, et al. The medaka draft genome and insights into vertebrate genome evolution. Nature. 2007;447(7145):714–719, doi:10.1038/nature05846.

40. Deguchi T, Itoh M, Urawa H, et al. Infrared laser-mediated retinal regeneration. Invest Ophthalmol Vis Sci. 2012;53(8):4810–4817, doi:10.1167/iovs.12-9511.

41. Ansai S, Kinoshita M. Targeted mutagenesis using local gene induction in medaka, zebrafish and Arabidopsis thaliana. Dev Growth Differ. 2019;61(1):114–140, doi:10.1111/dgd.12766.

42. Goto M, Harada Y, et al. Chromosome painting and systematic manipulation of morphological and motion cues. Cell Biol Open. 2014;3(5):362–371, doi:10.1242/bio.2014.04.021.

43. Dong S, Kang M, Wu X, Ye T. Development of a promiscuous fish model (Oryzias melastigma) for assessing multiple responses to stresses in the marine environment. Biomed Res Int. 2014;2014:563131, doi:10.1155/2014/563131.

44. Matsuo M, Kamey M, Fukamachi S. Behavioural red-light sensitivity in fish according to the otopotomor response. R Soc Open Sci. 2021;8(8):210415, doi:10.1098/rsos.210415.

45. Kayo D, Zempo B, Tomihara S, Oka Y, Kanda S. Gene knockout analysis reveals essentiality of estrogen receptor β1 (Esr2a) for female reproduction in medaka. Sci Rep. 2019;9(1):8868, doi:10.1038/s41598-019-45373-y.

46. Matsuo M, Ando Y, Kamey M, Fukamachi S. A semi-automatic and quantitative method to evaluate behavioral photosensitivity in animals based on the otopotomor response (OMR). Biol Open. 2018;7(6):bio035175, doi:10.1242/bio.035175.

47. Harada Y, Matsuo M, Kamey M, Goto M, Fukamachi S. Evolutionary history of the medaka long-wavelength sensitive genes and effects of artificial regression by gene loss on behavioural photosensitivity. Sci Rep. 2019;9:2726, doi:10.1038/s41598-019-39978-6.

48. Homma N, Harada Y, Uchikawa T, Kamey M, Fukamachi S. Protanopia (red color-blindness) in medaka: a simple model for producing color-blind fish and testing their spectral sensitivity. BMC Genet. 2017;18:10, doi:10.1186/s12863-017-0477-7.

49. Kamijo M, Kawamura M, Fukamachi S. Loss of red opsin genes relaxes sexual isolation between skin-colour variants of medaka. Behav Processes. 2018;150:25–28, doi:10.1016/j.beproc.2018.02.006.

50. Novales Flamarique I. Diminished foraging performance of a mutant zebrafish with reduced population of ultraviolet cones. Proc R Soc B Biol Sci. 2016;283(1826):2160058, doi:10.1098/rspb.2016.0058.

51. Flamarique IN. Opsin switch reveals function of the ultraviolet cone in fish foraging. Proc R Soc B Biol Sci. 2013;280(1752):20122490, doi:10.1098/rspb.2012.2490.

52. Daniele LL, Insinna C, Chance B, Wang J, Nikonov SS, Pugh EN. A mouse M-opsin monochromat: retinal cone photoreceptors have increased M-opsin expression when S-opsin is knocked out. Vision Res. 2011;51(4):447–458, doi:10.1016/j.visres.2010.12.017.

53. Greenwald SH, Kuchenbecker JA, Roberson DK, Neitz M, Neitz J. S-opsin knockout mice with the endogenous M-opsin gene replaced by an L-opsin variant. Vis Neurosci. 2014;31(1):25–37, doi:10.1097/01.wvn.2020.00801.

54. Novales Flamarique I, Fujihara R, Yazawa R, Bolstad K, Gowen B, Yoshizaki G. Disrupted eye and head development in rainbow trout with reduced ultraviolet (sws1) opsin expression. J Comp Neurol. 2021;529(11):3013–3031, doi:10.1002/cne.25144.

55. Kanazawa N, Goto M, Harada Y, et al. Changes in a cone opsin repertoire affect color-dependent social behavior in medaka but not behavioral photosensitivity. Front Genet. 2020;11:801, doi:10.3389/fgen.2020.00801.

56. Matsumoto Y, Fukamachi S, Mitani H, Kawamura S. Functional characterization of visual opsin repertoire in Medaka (Oryzias latipes). Gene. 2006;371(2):268–278, doi:10.1016/j.gene.2005.12.005.

57. Abio Madeira F, Mi Park Y, Lee J, et al. The EMBL-EBI search and sequence analysis tools APIs in 2019. Nucleic Acids Res. 2019;47(W3):W63–W64, doi:10.1093/nar/gkz268.

58. Sado Y, Inoue S, Tomono Y, Omori H. Advances in the identification and cloning of X-linked lymphocytes from enlarged iliac lymph nodes as fusion partners for the production of monoclonal antibodies after a single tail base immunization. Acta Histochem Cytochem. 2006;39(3):89–94, doi:10.1267/ahc.06001.

59. Kobayashi T, Namba M, Kohno M, Koyano T, Sado Y, Matsuyama M. An improved iliac lymph node method for production of monoclonal antibodies. Dev Growth Differ. 2022;64(1):38–47, doi:10.1111/dgd.12766.

60. Kusakabe T, Suzuki N. Photoreceptors and olfactory cells express the same retinal guanylyl cyclase isoform in medaka: visualization by promoter transgenesis. PLoS Lett. 2000;483(2–3):143–148, doi:10.1016/S0896-6273(00)80044-6.

61. Miyoshi K, Richards LJ, Akazawa C, O’Leary DDM, Nakashishi S. Labeling neural cells using adenoviral gene transfer into the brain: visualization by promoter transgenesis. Proc Natl Acad Sci U S A. 2004;101(8):2662–2667, doi:10.1073/pnas.0307534101.

62. Liu Y, Fisher DA, Storm DR. Intracellular sorting of neuro-modulin (GAP-43) mutants modified in the membrane targeting domain. 1994;14(10):8587–8517, doi:10.1523/JNEUROSCI.14-05-0807.1994.

63. Yoshida R, Sakurai D, Horie T, Kawakami I, Tsuda M, Kusakabe T. Identification of neuron-specific promoters in zebrafish intestinal. Genesis. 2004;39(2):130–140, doi:10.1002/gene.20032.

64. Moriyski K, Richards LJ, Akazawa C, O’Leary DDM, Nakashishi S. Labeling neural cells using adenoviral gene transfer into the brain: visualization by promoter transgenesis. Proc Natl Acad Sci U S A. 2004;101(8):2662–2667, doi:10.1073/pnas.0307534101.

65. Kusakabe R, Kusakabe T, Suzuki N. In vivo analysis of two striated muscle actin promoters reveals combinations of multiple regulatory modules required for skeletal and cardiac muscle-specific gene expression. Int J Dev Biol. 1999;43(6):541–554, doi:10.1387/ijdb.990610027.

66. Salbreux G, Barthel LK, Raymond PA, Lubensky DK. Coupling mechanical deformations and planar cell polarity to create regular patterns in the zebrafish retina. PLoS Comput Biol. 2012;8(8):1002618, doi:10.1371/journal.pcbi.1002618.
67. Schindelin J, Arganda-Carreras I, Frise E, et al. Fiji: an open-source platform for biological-image analysis. Nat Methods. 2012;9(7):676–682, doi:10.1038/nmeth.1929.
68. Schmid B, Schindelin J, Cardona A, Longair M, Heisenberg M. A high-level 3D visualization API for Java and ImageJ. BMC Bioinform. 2010;11:274, doi:10.1186/1471-2105-11-274.
69. Palczewski K, Kumakura T, Hori T, et al. Crystal structure of rhodopsin: a G protein-coupled receptor. Science. 2000;289(5480):739–745, doi:10.1126/science.289.5480.739.
70. Watanabe K, Nishimura Y, Oka T, et al. In vivo imaging of zebrafish retinal cells using fluorescent coumarin derivatives. BMC Neurosci. 2010;11:116, doi:10.1186/1471-2202-11-116.
71. Molday RS, MacKenzie D. Monoclonal antibodies to rhodopsin: characterization, cross-reactivity, and application as structural probes. Biochemistry. 1983;22(3):653–660, doi:10.1021/bi00272a020.
72. Ogawa Y, Shiraki T, Kojima D, Fukada Y. Homeobox transcription factor Six7 governs expression of green opsin genes in zebrafish. Proc Biol Sci. 2015;282(1812):21050659, doi:10.1098/rspb.2015.0659.
73. MacDonald IM, Tran M, Musarella MA. Ocular genetics: current understanding. Surv Ophthalmol. 2004;49(2):159–196, doi:10.1016/j.survophthal.2003.12.003.
74. Neitz J, Neitz M. The genetics of normal and defective color vision. Vision Res. 2011;51(7):653–651, doi:10.1016/j.visres.2010.12.002.
75. Sullivan LS, Bowne SJ, Birch DG, et al. Prevalence of disease-causing mutations in families with autosomal dominant retinitis pigmentosa: a screen of known genes in 200 families. Invest Ophthalmol Vis Sci. 2006;47(7):3052–3064, doi:10.1167/iovs.05-1443.
76. Rivas MA, Vecino E. Animal models and different therapies for treatment of retinitis pigmentosa. Histol Histopathol. 2009;24(10):1295–1322, doi:10.14470/hh-24.1295.
77. Jaissele GB, May CA, Reinhard J, et al. Evaluation of file rhodopsin knockout mouse as a model of pure cone function. Invest Ophthalmol Vis Sci. 2001;42(2):503–513.
78. Zelinka CP, Sotolongo-Lopez M, Fadool JM. Targeted disruption of the endogenous zebrafish rhodopsin locus as models of rapid rod photoreceptor degeneration. Mol Vis. 2018;24:587–602.
79. Zhang Y, Deng WT, Du W, et al. Gene-based therapy in a mouse model of blue cone monochromacy. Sci Rep. 2017;7(1):6690, doi:10.1038/s41598-017-06982-7.
80. Deng WT, Li J, Zhu P, et al. Rescue of M-cone function in aged Opn1mw-/- mice, a model for late-stage blue cone monochromacy. Invest Ophthalmol Vis Sci. 2019;60(10):3644–3651, doi:10.1167/IOVS.19-27079.
81. Fraser B, DuVal MG, Wang H, Allison WT. Regeneration of cone photoreceptors when cell ablation is primarily restricted to a particular cone subtype. PLoS One. 2013;8(1):e55541, doi:10.1371/journal.pone.0055410.
82. Yoshimatsu T, D’Orazi FD, Gamlin CR, et al. Presynaptic partner selection during retinal circuit reassembly varies with timing of neuronal regeneration in vivo. Nat Commun. 2016;7:10590, doi:10.1038/ncomms10590.
83. D’Orazi FD, Suzuki SC, Darling N, Wong RO, Yoshimatsu T. Conditional and biased regeneration of cone photoreceptor types in the zebrafish retina. J Comp Neurol. 2020;529(17):2816–2850, doi:10.1002/cne.24933.
84. Yoshimatsu T, Williams PR, D’Orazi FD, et al. Transmission from the dominant input shapes the stereotopic ratio of photoreceptor inputs onto horizontal cells. Nat Commun. 2014;5:3699, doi:10.1038/ncomms4699.
85. Rister J, Desplan C. The retinal mosaics of opsin expression in invertebrates and vertebrates. Dev Neurobiol. 2011;71(12):1212–1226, doi:10.1002/dneu.20905.
86. Raymond PA, Barthel LK, Curran GA. Developmental patterning of rod and cone photoreceptors in embryonic zebrafish. J Comp Neurol. 1995;359:537–550.
87. Cheng CL, Gan KJ, Flamarique IN. The ultraviolet opsin is the first opsin expressed during retinal development of salmonid fishes. Invest Ophthalmol Vis Sci. 2007;48(2):866–875, doi:10.1167/iovs.06-0442.
88. Krogh A, Larsson B, Von Heijne G, Sonnhammer ELL. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. J Mol Biol. 2001;305(3):567–580, doi:10.1006/jmbi.2000.4315.

**Supplementary Material**

**Supplementary Video.** The whole mounted retina of Tg(sws1:mem-egfp) medaka labeled with anti-GFP (green) and anti-SWS1 (pseudocolored violet) antibodies. The synaptic terminal of a cone cell appeared first, followed by the axon, the inner segment, and finally, the outer segment.