Orthologous Divergence and Paralogous Anticonvergence in Molecular Evolution of Triplicated Green Opsin Genes in Medaka Fish, Genus Oryzias

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Abstract

Gene duplication of green (RH2) opsin genes and their spectral differentiation are well documented in many teleost fish. However, their evolutionary divergence or conservation patterns among phylogenetically close but ecologically diverse species is not well explored. Medaka fish (genus Oryzias) are broadly distributed in fresh and brackish waters of Asia, with many species being laboratory-housed and feasible for genetic studies. We previously showed that a Japan strain (HNI) of medaka (Oryzias latipes) possessed three RH2 opsin genes (RH2-A, RH2-B, and RH2-C) encoding spectrally divergent photopigments. Here, we examined the three RH2 opsin genes from six Oryzias species representing three species groups: the latipes, the celebensis, and the javanicus. Photopigment reconstitution revealed that the peak absorption spectra ($i_{max}$) of RH2-A were divergent among the species (447–469 nm), whereas those of RH2-B and RH2-C were conservative (516–519 and 486–493 nm, respectively). For the RH2-A opsins, the largest spectral shift was detected in the phylogenetic branch leading to the latipes group. A single amino acid replacement T94C explained most of the spectral shift. For RH2-B and -C opsins, we detected tracts of gene conversion between the two genes homogenizing them. Nevertheless, several amino acid differences were maintained. We showed that the spectral difference between the two opsins was attributed to largely the E/Q amino acid difference at the site 122 and to several sites with individually small spectral effects. These results depict dynamism of spectral divergence of orthologous and paralogous green opsin genes in phylogenetically close but ecologically diverse species exemplified by medaka.

Key words: RH2 opsin, absorption spectra, spectral tuning, gene duplication, gene conversion, natural selection.

Introduction

Opsin genes responsible for color vision form a gene family created by gene duplications. Vertebrate cone opsins are classified into four phylogenetic types in vertebrates with different spectral sensitivities expressed in cone photoreceptor cells in the retina: SWS1 sensitive roughly to ultraviolet to violet, SWS2 to blue, RH2 to green, M/LWS to green to red ranges of the light wavelength (Yokoyama 2000). Teleost fish are unique among vertebrates in that the four types of opsin genes are often further duplicated to create diverse sets of subtypes (Chinen et al. 2003; Fuller et al. 2004; Parry et al. 2005; Matsumoto et al. 2006; Watson et al. 2011; Nakamura et al. 2013; Kasagi et al. 2015; Musilova et al. 2019). The reported subtypes are often differentiated in spectral sensitivity and spatiotemporal expression pattern in the retina presumably adapting to varying photic environments in water (Takechi and Kawamura 2005; Carleton et al. 2008; Kawamura et al. 2016; Mackin et al. 2019).

Among the four opsin types, the RH2 type appears to have experienced the largest number of gene duplication events (Spady et al. 2006; Rennison et al. 2012). The evolutionary rate of RH2 opsins in euteleost lineages is accelerated by 5- to 6-fold from that in the early vertebrate ancestors (Yokoyama and Tada 2010), implying that RH2 opsins of teleosts have
evolved in quite a dynamic fashion. Thus, RH2 opsins would be suitable for studying evolutionary fate and adaptation of duplicated opsin genes by examining their divergence or conservation patterns among phylogenetically close but ecologically diverse species, which is, however, not well explored.

The medaka (genus *Oryzias*), also known as ricefish, ranges broadly throughout fresh and brackish waters of Central, South, Southeast to East Asia, and the Indo-Malay-Philippines Archipelago as far east as Timor, and are currently classified into 24 species (Naruse 1996; Parenti 2008). *Oryzias* species can be divided into three major species groups, the *latipes*, *celebensis*, and *javanicus* groups (Takehana et al. 2005). Because of small size, relatively large and clear eggs, ease of maintenance in freshwater aquaria, and other reasons, the Japanese medaka (*Oryzias latipes*) has been one of the most widely used species in experimental biology and is one of the three most important model organisms among bony fish along with the zebrafish, *Danio rerio* and the pufferfish, *Takifugu rubripes* (Wittbrodt et al. 2002; Kasahara et al. 2007; Parenti 2008). Many *Oryzias* species are laboratory-housed and feasible for genetic studies (Iwamatsu et al. 1993; Matsuyama 1994; Katsumura et al. 2009; Matsumoto et al. 2009).

We previously examined a northern Japan strain, HNI, of *O. latipes* and showed that it possessed three RH2 opsin genes (*RH2-A, RH2-B, and RH2-C*) arrayed in tandem in the genome and encoding spectrally divergent photopigments with peak absorption spectra (*λ*-max) at 452, 516, and 492 nm, respectively (Matsumoto et al. 2006). Although the northern and southern populations of Japanese medaka are classified as the same species, being able to breed to produce healthy and fertile offspring, their split is estimated to be as deep as human–chimpanzee to human–orangutan splits, 4–18 Ma, and would have resulted in a genetic divergence with sufficient resolution for molecular evolutionary analyses (Takehana et al. 2003; Setiamarga et al. 2009). Thus, *Oryzias* species are suitable for the study of divergence or conservation patterns of the RH2 opsin gene among phylogenetically close (congeneric) but ecologically diverse species. In the present study, we examined the RH2 opsin genes of six species of *Oryzias* representing the three species groups: a southern Japan strain, HNIR, of *O. latipes* and Philippine medaka (*Oryzias luzonensis*) from the *latipes* group; Celebes ricefish (*Oryzias celebensis*) and Sarasins’ buntingi (*O. sarasinorum*) from the *celebensis* group; deep-bodied ricefish (*Oryzias melastigma*) and dwarf medaka (*Oryzias minutillus*) from the *javanicus* group (Formacion and Uwa 1985; Roberts 1998; Parenti and Soeroto 2004; Parenti 2008) (table 1).

## Materials and Methods

### Fish Samples

Five *Oryzias* species ( *O. luzonensis*, *O. celebensis*, *O. sarasinorum*, *O. melastigma*, and *O. minutillus*) were obtained from laboratory stocks supplied by NBRP Medaka (https://shigen.nig.ac.jp/medaka/, last accessed June 6, 2020) and maintained at the University of Tokyo. The University of Tokyo Animal Care and Use Committee approved all animal protocols (approval number C-14-01) in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki).

### DNA Cloning

Using the total RNA prepared from eyes of an adult fish, the first-strand cDNA was synthesized by using a poly (dT) primer (5’-aagcagtggtaacaacgcagagtact(30)vn-3’) (v: a, g, or c; n: a, g, c, or t; t(30): 30 succession of t). Opsin cDNAs were PCR-amplified using primer pairs (supplementary table S1, Supplementary Material online) which were designed based on the published nucleotide sequences of *RH2-A, RH2-B*, and *RH2-C* genes of *O. latipes* (HNI strain) (Matsumoto et al. 2006). Next, the 5′ and 3′ rapid amplification of cDNA ends (RACE) was carried out to reveal their untranslated regions and full-length coding sequences by using primer pairs specific to the species and genes (supplementary table S2, Supplementary Material online). The genomic sequences of the three genes including introns were determined for the same individuals with those examined for cDNA sequences using primers designed for photopigment reconstitution (supplementary table S3, Supplementary Material online). The PCR products were cloned into the pBluescript II (SK-) plasmids. All PCR products were sequenced in both strands using ABI PRISM 3130-Avant Genetic Analyzer (Applied Biosystems Japan, Tokyo, Japan). The DNA sequences were confirmed in duplicate PCRs.

### Phylogenetic Tree

Alignment of deduced amino acid sequences was carried out using CLUSTAL W (Thompson et al. 1994) and refined visually. Coding nucleotide sequences were aligned in accordance with the protein alignments. Evolutionary distance was estimated using the Tamura–Nei algorithm (Tamura and Nei 1993). Phylogenetic tree was reconstructed using the neighbor-joining method (Saitou and Nei 1987). The reliability of the tree topology was evaluated by the bootstrap analysis with 1,000 replications (Felsenstein 1985). All these analyses were conducted using a program package MEGA X (Kumar et al. 2018).

### Reconstitution of Opsin Photopigment

The opsin cDNAs were recloned into the pMT5 expression vector which contained the last 15 amino acids of the bovine rhodopsin necessary for immunoaffinity purification by 1D4 monoclonal antibody (Kawamura and Yokoyama 1998). The PCR primer pairs (supplementary table S3, Supplementary Material online) for the recloning contain the 5′- and 3′-edges
of the coding regions with necessary restriction sites for cloning and the Kozak sequence for efficient translation (Kawamura and Yokoyama 1998). The nucleotide sequences of the pMT5-cDNA clones were confirmed to match those of the template pBluescript-cDNA clones.

Point mutations were introduced by using QuikChange site-directed mutation kit (Stratagene, La Jolla, CA). All mutagenized cDNAs were sequenced in both strands using ABI PRISM 3130-Avant Genetic Analyzer (Applied Biosystems Japan, Tokyo, Japan).

The pMT5-cDNA clones were expressed in cultured COS-1 cells (RIKEN Cell Bank, Tsukuba, Japan). The cells were incubated with 5 µM 11-cis retinal (Storm Eye Institute, Medical University of South Carolina, Charleston, SC) and solubilized with 1% dodecyl maltoside (Anatrace, Maumee, OH). Produced photopigments were purified using the immobilized 1D4 monoclonal antibody (Cell Culture Center, Minneapolis, MN) as in previous studies (Chinen et al. 2003; Matsumoto et al. 2006). UV-visible absorption spectra of the photopigments were measured using U3010 dual-beam spectrometer (Hitachi, Tokyo, Japan) at 20°C for at least five times in the dark and for at least five more times after 3 min of light exposure.

Inference of Ancestral Amino Acid Sequences of RH2-A Opsins

The ancestral amino acid sequence at each ancestral node was inferred by using phylogenetic analysis by maximum likelihood (PAML) computer program version 4.8 with a likelihood-based Bayesian method (http://abacus.gene.ucl.ac.uk/software/paml.html, last accessed June 6, 2020) (Yang et al. 2003; Matsumoto et al. 2006). UV-visible absorption spectra of the photopigments were measured using U3010 dual-beam spectrometer (Hitachi, Tokyo, Japan) at 20°C for at least five times in the dark and for at least five more times after 3 min of light exposure.

Table 1

| Species Group | Species Name      | Distribution                                      | Habitat                        |
|---------------|-------------------|---------------------------------------------------|--------------------------------|
| latipes       | Oryzias latipes   | Eastern China, east Korea, Japanese Archipelago    | Fresh to brackish water        |
| latipes       | Oryzias latipes   | Luzon Island                                      | Creeks and rice fields         |
| celebensis    | Oryzias celebensis| South-western arm of Sulawesi, Lake Tempe, East Timor, River Mota Talau area | Inland and coastal rivers and streams |
| javanicus     | Oryzias sarasinorum | Central Sulawesi, Lake Lindu                       | Pelagic open waters            |
| javanicus     | Oryzias melanogaster| Indonesia, Sri Lanka, Bangladesh, and Myanmar      | Coastal brackish and freshwater |
| javanicus     | Oryzias minitillus | Chao Phrya basin and Salween basins, Mekong basin in northern Thailand and Kampuchea, Yunnan Province of China | Clear water swamps             |
| javanicus     | Oryzias minutillus| Thailand and Kampuchea, Yunnan Province of China   |                                |

Inference of Positive Selection in RH2-A Opsin Genes

To explore selective constraint of RH2-A opsin genes, same phylogenetic trees with inference of ancestral amino acid sequence of RH2-A opsin genes were used. Branch-site model, which was performed by the codeml program of the PAML 4.8 software package, allows the ω ratio to vary among sites along branches on the tree, such that positive selection can be detected in specific codon sites along particular lineages of interest. For branch-site model, Model A test was applied (Yang 2007; Yang and Dos Reis 2011), which estimates ω0 between 0 and 1 (site under purifying selection) and ω1 = 1 (sites evolving neutrally) for sites along all branches, as well as ω2a (sites under positive selection in the foreground branch but under purifying selection on the background branch) and ω2b (sites under positive selection in the foreground branch, which are evolving neutrally on the background branch). The likelihood ratio tests were performed between the Model A and the null Model A, which is the same Model A but with ω2 = 1 fixed. Likelihood ratio tests were carried out by comparing twice the difference in In likelihood scores of nested models against the number of extra parameters estimated by the more complex model. The Bayes Empirical Bayes calculation of posterior probabilities for site classes was used to calculate the probabilities of sites under positive selection.

Results

Phylogeny of Oryzias RH2 Opsin Genes

The cDNA and genomic sequences were determined for RH2-A, RH2-B, and RH2-C of O. luzonensis, O. celebensis, O. sarasinorum, O. melanogaster, O. minutillus, and O. nigroviridis (see Genome Biol. Evol. 12(6):911–923 doi:10.1093/gbe/evaa111 Advance Access publication 28 May 2020 913
supplementary tables S1–S3, Supplementary Material online, for relevant PCR primers). GenBank accession numbers of these nucleotide sequences are listed in supplementary table S4, Supplementary Material online. Regarding *Hd-rR* strain of *O. latipes*, nucleotide sequences of the three genes were retrieved from the whole genome sequence data (Kasahara et al. 2007) (supplementary fig. S1, Supplementary Material online).

Figure 1 shows a phylogenetic relationship reconstructed using coding nucleotide sequences among the *Oryzias* RH2 opsin genes including those of HNI (Matsumoto et al. 2006) with outgroup sequences from cichlid and zebrafish. All of the RH2-A sequences determined in this study clustered with those of HNI and Hd-rR strains of *O. latipes* and with the cichlid ortholog RH2B located outside of the *Oryzias* RH2-As as expected from previous studies (Matsumoto et al. 2006; Spady et al. 2006). Similarly, all the RH2-B and RH2-C sequences determined in this study clustered with those of HNI and Hd-rR and with the cichlid RH2Az and RH2Aβ.

**Gene Conversion between RH2-B and RH2-C**

In figure 1, RH2-B and RH2-C genes clustered by species group or by species as if they arose in the common ancestor of *O. latipes* and *O. luzonensis* (latipes group), in that of *O. celebensis* and *O. sarasinorum* (celebensis group), in *O. melastigma* and in *O. minutillus* by independent gene duplications. To test if this topology of the gene tree is due to sequence homogenization by gene conversions between the two genes, we applied GENECONV (Sawyer 1989) to the two genes in each *Oryzias* species (fig. 2A and supplementary table S5, Supplementary Material online). GENECONV searches for a consecutively identical region between two sequences and evaluates the probability of achieving homogeneity when we assume nucleotide differences to distribute randomly throughout the sequence region. An orthologous region corresponding to the nucleotide position 300 to position 918 in RH2-B of the *O. latipes* (HNI) appeared less affected by gene conversion (fig. 2A). In the reconstructed gene tree with
Absorption Spectra of *Oryzias* RH2 Opsins

Absorption spectra were measured for *Oryzias* RH2 opsin photopigments ([supplementary table S6](#), [Supplementary Material](#) online and [fig. 3](#)). The spectra in the dark had a prominent absorption peak in addition to a protein absorbance at 280 nm. When the reconstituted pigments were exposed to light, a new absorption peak appeared at
Fig. 3.—Absorption spectra of the reconstituted Oryzias RH2 opsin photopigments measured in the dark. The dark–light difference spectra are given in insets. $\lambda_{\text{max}}$ values are taken from the dark spectra. Because the amino acid sequence of the RH2-B opsin of O. latipes (Hd-rR) was identical with that of O. latipes (HNI), the $\lambda_{\text{max}}$ value of Hd-rR RH2-B was taken from the published information on HNI RH2-B.
Amino acid sequences were identical between A3 and A5, between A4 and O. latipes (Hd-R), and between A6 and O. minutillus. We thus reconstituted photopigments for A1, A2, and A3 and measured their absorption spectra (supplementary table S7 and fig. S4, Supplementary Material online). A2 was created by introducing three amino acid changes (Y95I [valine to isoleucine at residue 95], L275I, V281A) into O. luzonensis RH2-A. A1 was created from A2 by introducing four amino acid changes (C94T, V100M, F203Y, F255I). A3 (= A5) was created by introducing two amino acid changes (Y198F, R248K) to O. celebensis RH2-A. We also created the JTT-version of A1 by introducing L49I and M100V to the Dayhoff-version of A1. The $\lambda_{\text{max}}$ of the JTT-version of A1 was 467.0 ± 0.5 nm. Since the difference from the Dayhoff-version was negligible (1.7 nm), only the Dayhoff-version is used hereafter.

The largest spectral shift occurred in Branch A from A1 to A2 (−15 nm) toward the latipes group (O. latipes and O. luzonensis) (fig. 4). In the latipes group, further short-wave shifts occurred in Branch C (−6 nm toward O. latipes) and in Branch D (−7 nm toward O. luzonensis). Oryzias latipes HNI shifted oppositely for long wave with −5 nm. Toward the celebensis group and the javanicus group, spectral shift was moderate: −6 nm from A1 to A3 (the common ancestor of the two species groups) in Branch B with little shift thereafter except in Branch F toward O. celebensis with +6 nm long-wave shift.

**Complex Spectral Effects of Amino Acid Changes in Oryzias RH2-A Opsins**

Based on figure 4, we tested effects of the amino acid substitutions on the spectral shift occurred in the evolutionary branches A–F by site-directed mutagenesis (table 2). In Branch A with −15-nm spectral shift, there were four amino acid changes: T94C, M100V, Y203F, and I255F. A previous study on the newt SWS2 opsin reported a large effective spectral shift of the site 94 (Takahashi and Ebrey 2003). Indeed, T94C resulted in a 20-nm shift (table 2). However, this is larger than the actual spectral shift in this branch (−15 nm). When T94C was combined with single or double other mutations, spectral shifts were still larger. Thus, the spectral shift in Branch A was realized by nonlinear interaction of the four mutations with T94C playing a major role.

In Branch B with −6-nm shift, there were six amino acid changes: I48V, L49I, L105M, I112M, I162M, and S165C (fig. 4). Although an exhaustive survey was not conducted testing all single mutations and their combinations, the spectral shift was attributable to an additive effect of L105M and I162M with I112M and S165C with −6 nm long-wave shift.
enough spectral shift with −7 to −8 nm to explain the −6-nm shift of the branch. The other two also showed minor effects. Thus, nonadditive and regressive interaction among the five changes operates in this branch.

In Branch D with −7-nm shift, there were three amino acid changes: I95V, I275L, and A281V (fig. 4). Two of them, I275L and A281V, individually resulted in an enough spectral shift with −7 and −6 nm, respectively, to explain the −7-nm shift of the branch. But, when these two were combined, spectral shift was only −2 nm. Though I95V showed a minor effect with −2-nm shift, it extinguished the large effect of I275L when combined (−1 nm). Again, nonlinear and regressive interaction among mutations also operates in this branch.

In Branch E with +4-nm shift, there was only one change: S202T (fig. 4). Thus, the spectral shift of the branch should be attributed to this single mutation.

In Branch F with +6-nm shift, there were two amino acid changes: F198Y and K248R (fig. 4). Both mutations caused individually only minor and short-wave shifts with −1 and −2 nm, respectively. Thus, when combined, the two mutations exert a large synergistic effect with opposite wavelength direction from the effect of individual mutation.

| Table 2 |
| Spectral Effects of Mutations at Each Branch |

| Pigment | \( \lambda_{\text{Max}} \pm \text{SE (nm)} \) | \( \Delta \lambda \) from Ancestor |
|---------|--------------------------------|--------------------------|
| **Branch A** |
| Ancestor 1_T94C | 449.0 ± 0.9 | −20 |
| Ancestor 1_M100V | 467.0 ± 0.6 | −2 |
| Ancestor 1_T94CM100V | 448.3 ± 0.7 | −22 |
| Ancestor 1_T94CY203F | 449.7 ± 1.0 | −19 |
| Ancestor 1_T94C255F | 451.0 ± 1.4 | −18 |
| Ancestor 1_T94CM100VY203F | 450.0 ± 1.5 | −19 |
| Ancestor 1_T94CM100VI255F | 452.0 ± 1.3 | −17 |
| Ancestor 1_T94CY203FI255F | 449.5 ± 1.5 | −19 |
| **Branch B** |
| Ancestor 1_L49I | 468.0 ± 0.3 | −1 |
| Ancestor 1_L105M | 466.3 ± 1.1 | −2 |
| Ancestor 1_I112M | 465.7 ± 1.5 | −3 |
| Ancestor 1_S165C | 468.0 ± 0.4 | −1 |
| Ancestor 1_I48VL49I | 466.3 ± 0.4 | −2 |
| Ancestor 1_L105MI112M | 463.5 ± 0.4 | −5 |
| Ancestor 1_I162MS165C | 467.8 ± 1.0 | −1 |
| **Branch C** |
| Ancestor 2_F24Y | 450.7 ± 0.5 | −3 |
| Ancestor 2_L42I | 447.0 ± 1.0 | −7 |
| Ancestor 2_I108L | 446.0 ± 1.1 | −8 |
| Ancestor 2_I119L | 452.0 ± 0.9 | −2 |
| Ancestor 2_M309L | 447.3 ± 1.8 | −7 |
| **Branch D** |
| Ancestor 2_I95V | 452.0 ± 1.1 | −2 |
| Ancestor 2_I275L | 447.0 ± 1.1 | −7 |
| Ancestor 2_A281V | 448.0 ± 1.0 | −6 |
| Ancestor 2_I95VI275L | 453.0 ± 0.7 | −1 |
| Ancestor 2_I95VIA281V | 446.0 ± 1.0 | −8 |
| Ancestor 2_I275LA281V | 452.0 ± 1.1 | −2 |
| **Branch F** |
| Ancestor 5_F198Y | 462.0 ± 0.9 | −1 |
| Ancestor 5_K248R | 461.0 ± 0.8 | −2 |

Test of Natural Selection on Spectral Divergence of RH2-A
To explore whether amino acid replacements at Branch A in figure 4 were driven by positive Darwinian selection for functional divergence, we implemented branch-site model in PAML, indicating that the T94C substitution at Branch A was shown to have evolved under positive selection (supplementary table S8, Supplementary Material online).

Search for Conserved Amino Acid Differences between RH2-B and RH2-C Opsins against Gene Conversion as Key Sites for Their Spectral Distinction
Although the RH2-B and RH2-C opsin genes are homogenized by gene conversion (figs. 1 and 2), spectral difference is maintained between the RH2-B (\( \lambda_{\text{Max}} \) at 516–519 nm) and RH2-C (486–494 nm) opsins (fig. 3 and supplementary table S6, Supplementary Material online). Among the seven Oryzias species/stains (fig. 5), we searched for amino acid sites where in RH2-B opsins residues are same among at least five sequences, and in RH2-C opsins residues are also same among at least five sequences but are different from the major residue in RH2-B opsins. There were such sites in the N-terminal and transmembrane (TM) regions (fig. 5). Since TM regions are in
**Fig. 5.**—Aligned amino acid sequences of the RH2-B and RH2-C opsins among Oryzias species. Identical amino acids to the *O. latipes* (HNI) RH2-B sequence are indicated by dots. Gap sites are indicated with dashes. The seven transmembrane (TM) domains are indicated by horizontal lines above the *O. latipes* (HNI) RH2-B sequence. The sites 95, 96, 122, 123, 205, 209, 213, and 270 are highlighted by boxes.
the immediate vicinity to the retinal, we focused on the sites in
TM: 95, 96, 122, 123, 205, 209, 213, and 270 (highlighted in
fig. 5).

Using *O. latipes* (HNI) RH2-B and RH2-C cDNAs as tem-
plates, effects of the amino acid difference at the eight sites
on their 24-nm spectral difference were tested by site-
directed mutagenesis (fig. 6A–C). When all eight sites were
replaced in the RH2-C template to the RH2-B types (L95I/
N96T/Q122E/I123V/M205I/C209V/C213F/A270G), \( k_{\text{max}} \)
shifted to 516.0 ± 0.2 nm which matched well with that of
RH2-B (516.1 ± 0.5 nm) (fig. 6B). Similarly, when all eight sites
were replaced in the RH2-B template to the RH2-C types
(L95/L/T96/N/E122Q/V123I/I205M/V209C/F213C/G270A), \( k_{\text{max}} \)
shifted to 493.4 ± 0.8 nm which matched well with that of
RH2-C (492.3 ± 0.9 nm) (fig. 6B). Thus, among 21 amino
acid differences between RH2-B and RH2-C opsins of
*O. latipes* (HNI), these eight amino acid differences are suffi-
cient to explain the spectral difference between them.

Among the eight amino acid sites, E/Q exchange at site
122 is reported to exert a large spectral shift of ~15 nm in
RH2 opsins of Comoran coelacanth (*Latimeria chalumnae*)
and zebrafish (Yokoyama et al. 1999; Chinen et al. 2005a).
Consistently at this site, E and Q are perfectly conserved in
RH2-B and RH2-C opsins, respectively (fig. 5). When Q122E
was introduced to the RH2-C template, +16 nm of spectral
shift was observed, explaining two-thirds of the spectral
difference (fig. 6C). In contrast, there was no or only small
additional spectral effects detected in other sites when muta-
tions were introduced individually together with Q122E
(fig. 6C) although we failed to reconstitute a measurable
photopigment for L95/Q122E.

At the site 123, two residues (V and I) are perfectly con-
served in RH2-B and RH2-C opsins, respectively, as in the ad-
jacent site 122 with a large spectral effect. When all but site
123 were replaced in the RH2-C template to the RH2-B types,
the \( k_{\text{max}} \) matched well with that of RH2-B (supplementary fig.
5A and B, Supplementary Material online), confirming no
effect of the site 123. Despite no spectral effect, the perfect
conservation at the site 123 could be due to a hitchhiking
effect by the close vicinity from the site 122. When other sites
than the 123 and one additional sites (other than the site 122)
were replaced in the RH2-C template to the RH2-B types, the
\( k_{\text{max}} \) approached well to that of RH2-B (supplementary fig.
5B, Supplementary Material online), supporting individually
small spectral effects of other sites than the site 122. In par-
ticular, no effect was detected for sites 205 and 270 (supple-
mentary fig. 5B, Supplementary Material online).

To confirm whether the site 205 has no spectral effect, we
replaced all but site 205 in the RH2-C template to the RH2-B
types. The \( k_{\text{max}} \) matched with that of RH2-B (supplementary
fig. 5C, Supplementary Material online), confirming that
the site 205 has no effect. Using the RH2-C as the template,
when other sites than the 123 and 205 sites together with one additional site were replaced to the RH2-B types, the $\lambda_{\text{max}}$ differed slightly from that of RH2-B although we failed to evaluate the effect of site 96 (supplementary fig. 5C, Supplementary Material online). This shows individually small spectral effects of these additional sites including site 270 of which spectral effect was not detectable in supplementary figure 5B, Supplementary Material online.

These results suggest that the spectral difference between RH2-B and RH2-C opsin genes in genus Oryzias is maintained largely by the amino acid difference at site 122 and additionally by collective effects from amino acid differences at sites including 95, 96, 209, 213, and 270 through natural selection against gene conversion.

**Discussion**

In the present study, we examined the green-sensitive RH2-A, RH2-B, and RH2-C opsin genes for orthologous and paralogous divergence at nucleotide sequence and at absorption spectrum of encoding photopigments among medaka fish, genus Oryzias, which are broadly distributed in fresh and brackish waters of Asia. Among six species studied, representing all three species groups latipes, celebensis, and javanicus, absorption spectra of RH2-A photopigments were divergent ($\lambda_{\text{max}}$ at 447–469 nm). Spectral divergence among the Oryzias RH2-A is caused largely (~15 nm) by a single amino acid replacement (T94C) and by collection of individually small effects. The T94C occurred in the phylogenetic branch leading to the latipes species group. This replacement appears to be driven by positive natural selection. Absorption spectra of RH2-B and RH2-C were distinct between the two opsins and conservative among species ($\lambda_{\text{max}}$ at 516–519 and 486–493 nm, respectively) despite that tracts of gene conversion were found to have homogenized the two genes. We showed that the spectral differentiation was realized largely by a single E/Q amino acid difference at the site 122 and by the collective effect from other sites with individually small spectral effects which are considered to be maintained by natural selection against the homogenization effect of gene conversion.

The amino acid site 122 is located in the transmembrane region surrounding the retinal and directly interacts with $\beta$-ionone ring of the 11-cis retinal (Ahuja et al. 2009). Amino acid replacements at the site 122 are known to cause large spectral shifts to RH1 (rhodopsins; dim-light-sensitive visual opsins produced in rod photoreceptor cells in the retina), RH2, and SWS2 opsin genes of diverse vertebrates (Sakmar et al. 1989; Yokoyama et al. 1999; Takahashi and Ebrey 2003; Chinen et al. 2005a). Similarly, the amino acid site 94 is in the close vicinity of the Schiff base linkage of the 11-cis retinal to K294 and its counterion E113 (Palczewski 2000). The spectral shift by T94C is probably via a direct electrostatic effect on the retinal chromophore (Takahashi and Ebrey 2003). The amino acid replacements S94A and C94A are reported to cause ~14- and ~16-nm shift to the SWS2 opsins of newt and bluefin killifish, respectively (Takahashi and Ebrey 2003; Yokoyama et al. 2007). However, when the S94A is introduced into the goldfish SWS2 opsin, it only causes ~3-nm shift (Chinen et al. 2005b), suggesting that the spectral effect at the site 94 depends largely on the background amino acid sequence.

Whereas spectral effects of amino acid replacements at the sites 94 and 122 were relatively clear, effects of remaining replacements were not in the three RH2 subtypes in this study. Collective effects from multiple amino acid changes are often not additive. Additivity, exemplified by the “five-sites” rule of the M/LWS opsins (Yokoyama et al. 2008), is rather exceptional. This makes prediction of $\lambda_{\text{max}}$ unreliable based on empirical association of reported amino acid compositions at particular amino acid sites and observed $\lambda_{\text{max}}$ values (Chinen et al. 2005a, 2005b; Matsumoto et al. 2014; Kawamura et al. 2016). Recently, prediction of $\lambda_{\text{max}}$ values was made based on the magnitude of selected angles in carbon bonds in the conformation of 11-cis retinal under various amino acid sequence environments with known $\lambda_{\text{max}}$ as measured within atomistic molecular simulations (Patel et al. 2018). Accuracy of the prediction can be improved as more and more examples of observed $\lambda_{\text{max}}$ values for amino acid sequences are accumulated. Our present data should be a valuable source for the improvement.

In zebrafish, expression of RH2-1 and RH2-2 opsin genes, of which $\lambda_{\text{max}}$ values are the shortest and the second shortest among the four RH2 opsins, initiates earlier than the longer wave subtypes (Takechi and Kawamura 2005). Expression of zebrafish LWS opsin genes is similar, with the shorter wavesensitive LWS-2 expressed earlier than LWS-1 (Takechi and Kawamura 2005). This chromatic organization of the larval zebrafish retina appears to fit well with the behavioral requirements for feeding on zooplankton in a shallow water column which shows enhanced contrast under the rich shorter wavelength of light in the natural environment (Novales Flamarique 2016; Zimmermann et al. 2018). Cichlids also express the different RH2 opsin genes depending on their growth stage and ambient light (Parry et al. 2005; Spady et al. 2006). The quantitative-PCR analysis for the adult retina showed that in O. latipes, the RH2-A and RH2-B were expressed at a higher level than RH2-C (Matsumoto et al. 2006). In comparison to zebrafish and cichlids, data for medaka are wanting on the spatiotemporal expression of these genes in the retina as well as difference of feeding ecology among species and among species groups at larval to adult stages. Our finding of the large variation of the shortest-wave-sensitive RH2-A among the RH2 subtypes should facilitate relevant studies at organismal and ecological levels. $\lambda_{\text{max}}$ values of RH2-B and RH2-C opsin genes were found to be strictly conserved against homogenization effect by the gene conversion between their genes. The gene conversion and the
spectral conservation are also reported for primate LWS and MWS opsin (Balding et al. 1992; Hiwatashi et al. 2011). In primates, this spectral separation maintains the trichromatic color vision (Hiwatashi et al. 2011). However, it remains to be elucidated if the maintenance of spectral separation between the RH2-B and RH2-C opsin in medaka is for maintenance of color vision in which the two opsin is involved.

The effect of gene conversion on the divergence between duplicated genes has been well studied. A theoretical model demonstrates that the process consists of three phases I–III. Phase I is the time during which divergence reaches its equilibrium value. In phase II, divergence fluctuates around the equilibrium, and divergence increases again in phase III (Teshima and Innan 2004). The gene conversion was not observed between the RH2-A and the RH2-B/C genes. Gene duplication that derived the RH2-A gene and the ancestral RH2-B/C gene is ancient, predating speciation among Acanthopterygii (~140 Ma) (Matsumoto et al. 2006; Betancur-R et al. 2017; Hughes et al. 2018). Thus, it is probable that the divergence between RH2-A and RH2-B/C has already been in the phase III.

Our analysis for the RH2 opsins of six medaka species (O. latipes, O. luzonensis, O. celebensis, O. sarasinorum, O. minitillus, and O. melastigma) implied that all three sub-type genes were retained by natural selection among the genus medaka and were functionally important. This is the first case that opsin among closely related congeneric species are reconstituted in vitro and compared for their absorption spectra. Medaka is an excellent model for examining the relationship between vision and photonic environments because of their diversity of photonic environment and feasibility of laboratory studies. In future work, more detailed analyses of spatio-temporal gene expression in the retina are required for opsin genes of congeneric medaka species. Such studies would further clarify how fishes use multiple opsin genes formed by gene duplications and have adapted to their photonic environments during evolution.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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**Author Contributions**

S.K. designed research. Y.M. performed research and analyzed data. H.M. and S.O. contributed research materials. Y.M. and S.K. wrote the article.

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