Pleiotropic functions of chordin gene causing drastic morphological changes in ornamental goldfish

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Breeders and fanciers have established many peculiar morphological phenotypes in ornamental goldfish. Among them, the twin-tail and dorsal-finless phenotypes have particularly intrigued early and recent researchers, as equivalent morphologies are extremely rare in nature. These two mutated phenotypes appeared almost simultaneously within a short time frame and were fixed in several strains. However, little is known about how these two different mutations could have co-occurred during such a short time period. Here, we demonstrate that the chordin gene, a key factor in dorsal-ventral patterning, is responsible not only for the twin-tail phenotype but also for the dorsal-finless phenotype. Our F2 backcrossing and functional analyses revealed that the penetrance/expressivity of the dorsal-finless phenotype can be suppressed by the wild-type allele of chdS. Based on these findings, we propose that chdSwt may have masked the expression of the dorsal-finless phenotype, acting as a capacitor buffering gene to allow accumulation of genetic mutations. Once this gene lost its original function in the twin-tail goldfish lineages, the dorsal-finless phenotype could be highly expressed. Thus, this study experimentally demonstrates that the rapid genetic fixation of morphological mutations during a short domestication time period may be related to the robustness of embryonic developmental mechanisms.

Studying the peculiar phenotypes of ornamental animals may provide insights into how animal body shapes are able to change over time. Although domestication processes occur over time scales that are relatively short compared to those required for natural large-scale morphological evolution, breeders and fanciers of dogs, birds and fish have succeeded in establishing many ornamental domesticated animals with highly diverged phenotypes. Among these ornamental animals, goldfish (Carassius auratus) exhibit especially highly diverged morphological variations in the skeletal system, which are extremely uncommon in naturally selected vertebrate species. For instance, the dorsal-finless goldfish strains (e.g., Ranchu) exhibit bifurcated caudal fins and lack dorsal fins, which have intrigued early and recent researchers due to the rareness of equivalent mutations in other vertebrate lineages. While a bifurcated caudal axial skeleton has not been identified in any vertebrate taxa, this morphology is genetically fixed in the twin-tail ornamental goldfish strain. The dorsal-finless mutation is also genetically fixed in a domesticated goldfish population as an ornamental variation, but equivalent intra-species variations are not found in teleost species. Based on known molecular phylogenetic relationships and surveys of Chinese archives and paintings, these two mutant phenotypes are thought to have quasi-simultaneously appeared and became genetically fixed in the dorsal-finless goldfish strain during a domestication process that lasted less than 600 years. It is also revealed that the chdS locus (or chdA locus in previous study)—the homologue of chordin gene which play a significant role for the dorsal-ventral patterning—was identified as the responsible gene of the twin-tail phenotype; a stop codon-containing allele (chdSE127X) is the responsible allele and the chdSE127X/E127X genotype is commonly shared among all of the investigated ornamental goldfish strains with bifurcated caudal fin, including Ryukin, Oranda, and Ranchi. Subsequently, genome-wide association study (GWAS) methodologies were used to identify multiple candidate loci that might be related with the dorsal-finless phenotype. However, little is known about how breeders and fanciers so quickly succeeded to genetically fixing the genes that give rise to such a peculiar ornamental goldfish strain, especially in light of the fact that the morphological mutations occur in multiple complex and sophisticated body parts.

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Here, we conducted genetic and functional analyses in the dorsal-finless phenotype focusing on the chdS locus. Since all of the investigated modern dorsal-finless goldfish strains share the twin-tail phenotype and chdS mutated genotype (chdS$^{E127X/E127X}$), it is reasonable to suspect that the chdS locus may be related not only with the bifurcated caudal fin but also with the dorsal-finless phenotype$^{6,7,11}$. To examine this possibility, we conducted genotyping and phenotyping of backcrossed F2 progenies derived from the hybrids of the single-tail common goldfish (wild-type) and the Ranchu strain (Fig. 1a–d)$^{6,11}$. Moreover, we performed microinjection of chdS$^{wt}$ mRNA into the progenies derived from Ranchu strain parents (Fig. 1e). Based on our results, we conclude that the chdS gene is indeed responsible for both the twin-tail and dorsal-finless phenotypes, suggesting that these two phenotypes are causally related to each other. These findings provide intriguing insights into how a highly conserved body architecture can be drastically changed during a short time period.

Results

Genetic backcross analysis. To conduct our genetic analysis, we generated F2 backcrossed progenies starting with F0 parents of the wild-type and Ranchu strain (Fig. 1d; Supplementary Fig. S1). The F0 wild-type and Ranchu parents were respectively chdS$^{wt/wt}$ and chdS$^{E127X/E127X}$ in their chdS genotypes (Fig. 1a,b). This set of F0 wild-type and Ranchu individuals yielded a total of 123 F1 progenies in four clutches: 2021-0426-22-RAwt (n = 43), 2018-0328-05 (n = 22), 2021-0510-09 (n = 24), 2021-0510-12 (n = 34). All of these F1 progenies carried a chdS$^{wt/E127X}$ genotype and exhibited single median fins (including dorsal, anal and caudal fins), similar to the wild-type goldfish (Fig. 1c). Based on this consistent phenotype, we can conclude that the alleles conferred by wild-type goldfish suppress the dorsal-finless phenotype in the F1 progenies.

The F1 progenies were then backcrossed with the Ranchu strain to generate F2 progenies (Fig. 1d, Supplementary Fig. S1). Two clutches of F2 progenies were obtained and designated “2020-0511-06RA” and “2021-0406-01RA”. From 116 individuals in these two clutches of F2 progenies, we successfully obtained phenotype and genotype data. Notably, the F2 progenies exhibited variations in their caudal and dorsal fins (Fig. 2a–h). For each F2 individual, either a single or bifurcated caudal fin was observed, consistent with our previous report$^{11}$. The dorsal-finless phenotypes were also varied, as shown in Fig. 2a–h. Most of the F2 progenies either completely lacked dorsal fin rays, similar to the Ranchu strain (Fig. 1b), or the animals had a wild-type-like dorsal fin (Fig. 2a,b,g,h). Nevertheless, some intermediate phenotypes were observed (Fig. 2c–f). For example, some individuals showed a rudimentary dorsal fin, while others had only a few dorsal fin rays (Fig. 2c–f).

To examine the influence of chdS$^{wt}$ and chdS$^{E127X}$ alleles on the dorsal fin phenotype, we counted the numbers of dorsal fin rays in the F2 backcross population and genotyped the chdS locus in 116 progenies (Fig. 2i,j; Supplementary Tables S1 and S2); the two clutches respectively contributed 67 and 49 individuals to the total. Our analyses of these F2 progenies clearly indicated that the chdS$^{wt}$ allele influences the dorsal-finless phenotype (Fig. 2a–f, Supplementary Tables S1 and S2). In both clutches, most of the individuals with a chdS$^{E127X/E127X}$ genotype exhibited bifurcated anal and caudal fins, although many still exhibited single anal and caudal fins (Supplementary Tables S3 and S4). These findings were again consistent with our previous report$^{11}$.

The chdS$^{E127X/E127X}$ progenies could be categorized as having wild-type and mutant dorsal fins based on the number of the dorsal fin rays present. Our previous research showed that most wild-type goldfish have a total

![Figure 1. Schematic views of experimental procedures. Lateral views of the wild-type (a) and Ranchu strain (b) goldfish. (c) A lateral view of hybrid progeny of Ranchu and wild-type. (d) Schematic representation of backcross analysis. (e) Schematic representation of mRNA microinjection rescue experiment. The size of the individuals in (a, b, and c) are 5 cm, 6 cm and 6 cm approximately, respectively.](https://doi.org/10.1038/s41598-022-24444-7)
number of dorsal fin rays ranging from 17 to 22 at the late larval stage (Pr stage), so we assigned a mutant phenotype classification to individuals with less than 17 dorsal fin rays in their dorsal fin. Based on this criterion, more than half of the chdSE127X/E127X progenies exhibited a dorsal fin mutation phenotype (Fig. 2i, j; Supplementary Tables S1–S4). In contrast to the chdSE127X/E127X individuals, the chdSwt/E127X F2 progenies tended to show the wild-type phenotype in the dorsal morphology (Fig. 2i, j; Supplementary Tables S1–S4). These results suggested that the presence of chdSwt significantly suppresses the expression of the dorsal-finless phenotype; nevertheless, several chdSwt/E127X F2 progenies showed anomalous mutated phenotypes at the dorsal fin (Fig. 2i, j; Supplementary Fig. 2a–d). In short, our data strongly imply that the expression of mutations in the median fins are genetically linked with chdS locus.

mRNA microinjection rescue. To definitively test whether the chdS locus influences the expression of the mutation phenotype in the dorsal fin and the other median fins, we designed a rescue experiment in Ranchu strain progenies. In this experiment, we performed chdS mRNA microinjection, which could rescue the twin-tail phenotype in our previous study. Microinjections of chdSwt mRNA were made into the fertilized eggs derived from Ranchu parents (Methods) (Fig. 1e). The experiment was repeated four times in independent clutches (#2022-0315-01RARA, #2022-0315-04RARA, #2022-0321-01RARA, #2022-0321-03RARA), allowing us to examine the phenotypes in a total of 1474 hatched larvae at 3 days post-fertilization (dpf) (Supplementary Table S5). The examinations focused on the morphologies of the median fin fold and its derivatives (Fig. 3a–n; Supplementary Fig. S2). Our rescue experiments demonstrated that chdSwt mRNA microinjected into early-stage larvae caused the expression of different phenotypes than those seen in the control larvae (non-injected or water-injected larvae), in terms of the median fin fold at pre- and post-cloacal levels (Fig. 3a–h). Moreover, these phenotypic variations of

![Figure 2. Dorsal fin phenotype in F2 hybrid. (a–h) Lateral views of the dorsal region [three representative variations of dorsal-finless mutants with bifurcated caudal fins (a–f) and wildtype (g, h)]. Numbers in the lower left corner indicate the number of dorsal fin rays including serrated dorsal fin spine and post-serrated dorsal fin spine rays (please see text in details). Panels of (b), (d), (f), and (h) are magnified views of (a, c, e, and g), respectively. (i, j) Histograms of distribution patterns of dorsal fin ray number of chdSwt/E127X and chdSE127X/E127X genotypes. (i) 2020–0511-06RAWT clutch, (j) 2021–0406-01-RAWT clutch). Green-colored dotted vertical line indicates the 17 dorsal fin ray number (the boundary of wild-type and mutant phenotype; see text).]
the median fin fold were related to the dorsal fin morphology at the late larval stage (Fig. 3i–n). All of the control hatched larvae exhibited a relatively uniform ventralized phenotype (e.g., bifurcated and/or disrupted fin folds at the ventral side; white asterisks, malformed fin fold at the ventral side; black asterisks, malformation of the anterior ventral side tissues). Panels of (b, d, f and h) are magnified views of (a, c, e, and g), respectively. Hatched larva exhibiting group 1 (a, b), group 2 (c, d) or group 3 (e–h) phenotypes. (i–n) Late larvae. Control (i, j) and chdS<sup>wt</sup> mRNA injected late larvae (k–n). Panels of (i, l) and (m) are magnified views of (i, k), and (m), respectively (black arrowhead, partially rescued dorsal-fin; white bracket, completely rescued dorsal-fin). Late larvae on panels on (i, k), and (m) are derived from the hatched larvae of group 1, group 2, and group 3, respectively. (o–r) Proportion of rescued specimens following injection of embryos with the wild type chdS mRNA (chi-square test p < 0.0001 in all clutches). Scale bars = 0.1 mm (h), 1 mm (g, m, n). Panels of hatched larvae (a, c, e, g), the magnified views of the hatched larvae (b, d, f, h), late larvae (i, k, m), and the magnified view of the late larvae (j, l, n) are shown at the same magnification.

Figure 3. Rescue of the dorsal-finless phenotype by mRNA microinjection. (a–h) Representative phenotypes of hatched larvae (3dpf). Control (a, b) or chdS<sup>wt</sup> mRNA injected hatched larvae (c–h) (arrowheads, partially rescued dorsal fin fold; black bracket, completely rescued fin fold; black asterisks, mutated fin fold at the ventral side; white asterisks, malformation of the anterior ventral side tissues). Panels of (b, d, f and h) are magnified views of (a, c, e, and g), respectively. Hatched larva exhibiting group 1 (a, b), group 2 (c, d) or group 3 (e–h) phenotypes. (i–n) Late larvae. Control (i, j) and chdS<sup>wt</sup> mRNA injected late larvae (k–n). Panels of (i, l) and (m) are magnified views of (i, k), and (m), respectively (black arrowhead, partially rescued dorsal-fin; white bracket, completely rescued dorsal-fin). Late larvae on panels on (i, k), and (m) are derived from the hatched larvae of group 1, group 2, and group 3, respectively. (o–r) Proportion of rescued specimens following injection of embryos with the wild type chdS mRNA (chi-square test p < 0.0001 in all clutches). Scale bars = 0.1 mm (h), 1 mm (g, m, n). Panels of hatched larvae (a, c, e, g), the magnified views of the hatched larvae (b, d, f, h), late larvae (i, k, m), and the magnified view of the late larvae (j, l, n) are shown at the same magnification.
Notably, the level of rescue in the hatched larvae appeared to vary in terms of the dorsal fin fold phenotype (Fig. 3c–h). Based on the dorsal fin morphology, we categorized the control and mRNA-injected larvae into three different groups as follows: (i) no dorsal fin fold at the level of yolk and post-yolk levels (group 1; Fig. 3a,b), (ii) partial dorsal fin fold at the level of yolk and post-yolk levels (group 2; Fig. 3c,d), and (iii) a wild-type equivalent phenotype of complete dorsal fin fold at the yolk and post-yolk levels (group 3; Fig. 3e–h) (for further descriptions of each group, see Methods). The morphological observations of these categorized larvae at the late larval stage (Fig. 3i–n) were entirely consistent with the early larval phenotypes (Fig. 3a–h). More specifically, at the late larval stages, dorsal fin rays were not observed in group 1 individuals, dorsal fin rays were missing at several body levels in group 2 individuals, and dorsal fin rays were completely developed in the group 3 individuals (Fig. 3i–n). This concordance of phenotypes suggested that chdS

**Discussion**

In this study, we show that the chdS gene is responsible not only for the twin-tail phenotype but also for the dorsal-finless phenotype, suggesting that the chdS gene has pleiotropic functions that influence the formation of mutated morphologies in all median fins. Based on our results, we posit an evolutionary process of the dorsal-finless phenotype. (1) Before the genetic fixation of the twin-tail phenotype, the penetrance/expressivity of the dorsal-finless phenotype was suppressed due to the chdS

Figure 4. Schematic representation of the appearance of the dorsal-finless phenotype. The topology of the phylogenetic tree is based on the molecular phylogenetic analyses. The magenta circle indicates the genetic fixation of the chdS

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loci in terms of phenotype expression. The fact that several F2 progenies with \( chdS^{wt/E127X} \) genotype expressed the dorsal-finless phenotype indicates that there is at least one other responsible locus for the dorsal-finless phenotype independent of the \( chdS \) gene (Fig. 2i, Supplementary Fig. S2a–d). However, our results from the F2 segregant analyses indicated that the majority of mutated alleles responsible for the dorsal-finless phenotype are under the influence of \( chdS \) gene. Thus, the \( chdS \) gene appears to be epistatic to most genes responsible for the dorsal-finless phenotype. Taking into consideration the difficulty of detecting epistatic effects in complex human phenotypes using GWAS studies as well as the complicated molecular networks of \( chordin \)-related genes, it may be extraordinarily difficult to unambiguously identify the genes responsible for the dorsal-finless phenotype with simple application of a GWAS design on the domesticated population of goldfish\(^{6,7,23} \).

In fact, there is still no consensus about the responsible genes for the dorsal-finless phenotype, even though two GWAS reports have been published on the dorsal-finless phenotype by two independent research groups\(^{6,7} \). One group identified \( lrp6 \) gene as a candidate gene for the dorsal-finless phenotype, while the other group suggested a relationship exists between the mutated phenotype and the \( dhfr \) gene\(^{6,7} \). These inconsistent findings cannot be adequately explained at the level of the molecular mechanism due to a paucity of studies on these genes in goldfish. However, our study suggests that the inconsistent findings from these GWAS reports might be at least partially due to the complex epistatic relationships between the gene regulatory network and the \( chordin \) gene\(^{6,24–27} \). Our study also strongly indicates that further investigations will be required to determine how the early embryonic developmental process relates to the simultaneous occurrence of the twin-tail and dorsal-finless phenotypes\(^{25–27} \).

Careful consideration of the embryonic proximity of the dorsal and caudal fins and their modular relationship may provide a deeper understanding of how these two phenotypes may have simultaneously occurred\(^{1,2,6,16,21,29,30} \). Molecular developmental studies in zebrafish revealed that the mesenchymal cells of the dorsal- and caudal fins are derived from the different levels of somite, but the epithelial cells are connected to each other at the embryonic stage; the epithelial cells form the fin fold and its primordia\(^{31,32} \). Based on the close embryonic proximity of epithelial cells, it is naturally presumed that both dorsal- and caudal fins are under the influence of the same molecular mechanisms. In fact, this interpretation allows us to explain why the majority of our F2 backcross progenies consistently exhibited mutant phenotypes in dorsal, anal, and caudal fins (Supplementary Tables S1, S2). Moreover, previous reports on \( szl \)-depleted goldfish are consistent with this interpretation\(^{18,30} \). Depletion of \( szl \) gene expression in goldfish embryos leads to the dorsal-finless phenotype, which is related to the \( chordin \) gene\(^{37} \). Although the detailed molecular developmental mechanisms of the dorsal fin disappearance are still unknown, it is certain that the depletion of this gene disrupts dorsal–ventral patterning to generate the dorsal-finless phenotype in the goldfish\(^{33} \). Thus, it is reasonable to assume that the morphogenesis processes of the dorsal and caudal fin are under the regulation of the same dorsal–ventral patterning molecular developmental mechanism (which involves the \( chordin \) genes) at the early embryonic stage, even though these fins are separate structures at juvenile and adult stages.

We can rephrase the aforementioned assumption as follows. The \( chdS^{wt} \) allele might have a buffering capacitor function that allows the dorsal–ventral patterning mechanisms to accumulate genetic mutations\(^{25–27} \). This assumption allows us to answer why the twin-tail and the dorsal-finless phenotype might appear quasi-simultaneously during the goldfish domestication history. The accumulation of responsible mutations for the dorsal-finless phenotype might have been masked by the \( chdS^{wt} \) allele, and the effects of these mutations could have been unmasked specifically in the twin-tail goldfish lineages. Of course, our present study cannot completely reject the proposition that all of the responsible mutated alleles of the dorsal-finless phenotype had appeared after the fixation of \( chdS^{E127X} \) in the goldfish population. Nevertheless, an evolutionary scenario that includes a buffering capacitor function of the \( chdS \) gene seems more plausible than a scenario in which de novo genetic mutations for the dorsal-finless phenotype appeared in the lineage of the Ranchu group, as the first scenario could easily explain why the fixation process of the dorsal-finless phenotype in the twin-tail ornamental goldfish as so rapid (Fig. 4).

Researchers have investigated how masked genetic variations (or cryptic genetic variations) contribute to the expression of phenotypes upon genetic perturbation in several vertebrate species, including teleost species\(^{34–37} \). For example, the effects of chaperone protein HSP90 have been investigated in zebrafish and cavefish\(^{34,35} \). Moreover, the \( mef2ca \) gene was also examined in terms of its buffering function in the context of dermal skeleton morphogenesis\(^{36,37} \). Unlike these experimentally derived phenotypic variations, the phenotypic variations in goldfish were established through a process of artificial selection for the stabilization of visually classifiable strains (for example, \( Ryukin \) and \( Ranchu \) strains) after the fixation of the \( chdS^{E127X} \) allele\(^{41} \). Thus, it is reasonable to assume that the ornamental morphologies of established goldfish strains are the consequence of the canalization of morphogenesis\(^{36–41} \). More specifically, the stable expression of both the twin-tail and dorsal-finless phenotypes has been canalized in the lineage of the \( Ranchu \) strain, but not in \( Ryukin \) strain. This fact would imply that the various dorsal-finless phenotype-associated candidate genes identified by GWAS analyses might actually contribute to the phenotypic canalization, rather than acting as decisive factors for the dorsal-finless phenotype\(^{25–27} \). We hope that future studies may combine the GWAS approach with molecular developmental genetics methodologies to better reveal the relationships between drastic morphological changes, developmental robustness, and genetic variations in different ornamental goldfish strains.

**Methods**

**Goldfish strains.** Goldfish were purchased from an aquarium fish agency in Taiwan. Two strains were used in this research, the single tail common goldfish (wild-type) and the \( Ranchu \) strain. To avoid confusion about the definition of “wild-type,” which may arise from differences in goldfish nomenclature systems used by breeders and researchers, this study defined goldfish individuals with a slender body and a single fin as having a wild-type phenotype (Fig. 1a)\(^{15} \). The \( Ranchu \) strain individuals lack a dorsal fin and exhibit a bifurcated caudal fin.
(Fig. 1b). The F0 parents were genotyped by PCR and restriction enzyme digestion as described in a previous paper. The wild-type goldfish parents were homozygous for the chdS<sup>wt</sup> allele, and dorsal-finless goldfish were homozygous for the chdS<sup>SE127X</sup> allele. By crossing these parental strains, F1 individuals were obtained (Fig. 1c). These F1 individuals were subsequently used to obtain F2 segregants. The research was performed in accordance with internationally recognized guidelines and ARRIVE guidelines. Ethical approval was from the Institutional Animal Care & Utilization Committee of Academia Sinica, Taiwan (Protocol ID: 19-11-1351).

**Artificial fertilization.** The procedure for artificial fertilization was based on our previous report. In the spawning season (March to June), sperm were taken from males and preserved in Modified Kurokura’s extender 2 solution at 4 °C. Eggs were squeezed out from mature females into Teflon-coated dishes. Artificial fertilization was performed using dry methods. The fertilized eggs were spread onto 9-cm Petri dishes, containing tap water (approx. 24 °C).

**Genotyping.** The genotype of chdS locus was examined by PCR and restriction enzyme digestion as previously reported. PCR primers were designed to amplify the region containing both the chdS<sup>SE127X</sup> allele and the closely linked Aval restriction enzyme site. PCR fragments amplified by these specific primers were digested by Aval, and separated on 2% agarose gels. Genotypes were determined on the basis of the resulting band patterns.

**mRNA microinjection rescue experiment.** The pCS2 + plasmid vector containing the coding regions of the chdS<sup>wt</sup> sequence were used to synthesize mRNA for the microinjection rescue experiments. The sequence containing the coding region of chdS<sup>wt</sup> was amplified by PCR reaction, using a forward primer Sp6 (5′-ATT TAGGTGACACTATAGA-3′) and reverse primer M13R (5′-TCACACAGAAAAACGCTA TGAC-3′). The PCR product was cleaned with a GeneMark DNA Clean/Extraction kit before being used as the DNA template to synthesize capped mRNA. The mRNA synthesis was performed with the mMESSAGE mACHINE SP6 kit, according to the manufacturer's instructions (Ambion Inc.). Then, the synthesized mRNA transcripts were purified using a Monarch RNA Cleanup kit and resuspended in nuclease-free water. The microinjection mixture contained 1 μl Phenol Red (Sigma), 1 μl of 2 M KCl, 4.7 μl nuclease-free water, and 3.3 μl synthesized mRNA (300 ng/μl) in a total of 10 μl. Phenol red (Sigma) was used as an indicator at a final concentration of 0.05%. Injection needles were prepared from borosilicate glass filaments (Sutter instrument, BF100-50-10) using a micropipette puller (Model P-1000, Sutter Instrument). A microinjector (Eppendorf Femtojet; Eppendorf) was used to inject mRNA into the center of the yolk of fertilized eggs at the 1–2 cell stage, which were maintained on Petri dishes. Each embryo was injected with 2 μl of the mRNA mixture containing a total of 250 pg mRNA. The injected embryos were then incubated at 24 °C until phenotyping at 3 dpf. Four independent rescue experiments were performed by injecting chdS<sup>wt</sup> mRNA into the embryos derived from Ranchu strain parents. At 3 dpf, embryos were categorized into three groups (described below). These categorized 3 dpf larvae were maintained for more than one month and assessed for their phenotype at hatching. To examine the potential influence of the mechanical contact with the injection needle, nuclease-free water was injected into the embryos as a control. As expected, mechanical contact did not cause significant changes to the morphology of the larvae during the developmental process.

**Phenotype analysis.** The F2 segregants were phenotyped at both late embryonic (3 dpf) and post-embryonic stages. The embryos and larvae were observed under stereomicroscopy. The 3 dpf embryos were categorized into three groups (group 1, group 2 and group 3) based on the position of rescued fin fold shape and its relative position to yolk; the position of dorsal fin rays at the juvenile stage is known to be related with that of the cloaca. The criteria for these categories were as follows: group 1 individuals had no dorsal fin fold at the level of yolk; group 2 had dorsal fin fold partially at the level of yolk; and group 3 had a complete dorsal fin fold at the yolk and post-yolk levels. The phenotyping of late larvae and juveniles was conducted based on the numbers of dorsal fin rays. Maintenance of larvae followed previous reports. The progenies from Pelvic fin ray stage to juvenile stage were fixed with PFA (4 wt% of paraformaldehyde in 1 × PBS solution) overnight, washed in 70% ethanol, and stained with alizarin red solution (0.02% alizarin red in 70% ethanol). Alizarin red-stained specimens were washed with 70% ethanol to reduce background. Phentotypic observations in larvae and stained specimens were made under stereomicroscopy (SZX16 and SZ16, Olympus). Images were acquired using a stereo microscope system with digital microscope camera (SZX16 with DP80; Olympus). All data comparisons were made with the R statistical computing package of RStudio (Build 554, version 2022.07.1 + 554).

**Data availability**

The dataset supporting the conclusion of this article are included within the article (Supplementary Tables S1–S5).

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**Author contributions**

G.A., and K.G.O. conceptualized the project. C.W., I.L., and H.C. conducted artificial fertilization and fish maintenance. H.C.C. performed molecular experiments. K.G.O wrote original draft. H.C., A.G., and K.G.O. revised and edited manuscript. All authors reviewed the manuscript.
Competing interests
The authors declare no competing interests.

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