Title: Production of androgenetic diploid loach by cold-shock of eggs fertilized with diploid sperm

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Abstract
Diploid androgenotes were produced without egg irradiation in the loach, *Misgurnus anguillicaudatus*. Eggs of wild-type diploid females were fertilized with the diploid sperm of a neo-tetraploid male, and then cold-shock treated at 3.0 (range, ± 0.5) °C for 30 min just after fertilization to eliminate the female nucleus. After hatching, the ploidy status of hatched larvae was analyzed by flow cytometry, which revealed putative diploid androgenotes as well as larvae possessing other ploidies. Five independent microsatellite DNA markers were genotyped to confirm all-male inheritance of the resultant diploid larvae. The yield rate of diploid androgenetic larvae to total eggs used was 12.29 ± 3.25% in the cold-shock group, and 22.23 ± 13.42% in the UV-irradiated group, and their difference was not statistically significant (*P* > 0.05). No diploid androgenetic larvae were detected in the intact control group. To our knowledge, this is the first report demonstrating the successful induction of diploid androgenotes without egg irradiation in fish.

**Keywords**: cold-shock; androgenesis; *Misgurnus anguillicaudatus*

1. **Introduction**
Teleost eggs are normally ovulated at the metaphase of the second meiosis (M II). At this point, the eggs are physiologically mature and are spawned outside of the female to accept sperm for fertilization in ambient water. Such reproductive traits of most teleosts provide the potential for chromosome manipulation, such as induced polyploidy (increase in the number of chromosome sets), gynogenesis (all-female inheritance), and androgenesis (all-male inheritance). Androgenesis is also defined as the development of progeny carrying chromosomes exclusively transmitted from the male parent. Three types of androgenesis have been recognized to date: (1) obligate androgenesis (or paternal apomixis), (2) spontaneous androgenesis, and (3) artificial androgenesis [1]. In obligate androgenesis, all progeny inherit the paternally derived nuclear genome and reproduce without any contribution of the maternally derived nuclear genome. Although this type of androgenesis has been demonstrated in the bivalve species, *Corbicula* [2-6], and in arthropods [7], its frequency is very low in the animal kingdom overall. Spontaneous androgenesis also occurs at very low rates in gonochoristic species that reproduce bisexually [1]. With respect to teleost fish species, spontaneous androgenesis has been found to occur at a frequency of approximately 1% in an intergeneric hybrid between the common carp (*Cyprinus carpio*) and the grass carp (*Ctenopharyngodon idella*) [8-11], and was also very recently observed to occur at a frequency of approximately 5% in a cross between different clonal strains of the silver crucian carp (*Carassius auratus gibelio*) [12]. Since obligate and spontaneous androgenesis are generally rare events in teleost
fish and other aquatic animals, induction of artificial androgenesis has been extensively studied and several inducing protocols have been put forward [13-16]. All of the previous protocols involve the irradiation of eggs in the first step of the process in order to genetically inactivate the maternal genome (egg nucleus) before fertilization. Generally, gamma, X, or ultraviolet (UV) rays are used as the irradiation source. While these methods are effective for inactivating the egg nucleus, they are not particularly convenient for routine use [16]. The solutions used during irradiation, such as ovarian fluid [17], seminal plasma [18-19], Ringer’s solution [20], or Hank’s solution [21], are essentially required not only to protect the eggs from dryness, but also to dilute the density of eggs to enable uniform irradiation while maintaining fertilizing ability during the process. Moreover, in the case of gamma ray irradiation, the procedure must be performed in a special facility for safety purposes.

To overcome the limitations involved in the irradiation process, Morishima et al. [22] proposed a new method for the artificial induction of haploid androgenesis in the loach, *Misgurnus anguillicaudatus*, without the use of UV-irradiation. Their method involves cold-shocking eggs at a temperature of 0 to 3 °C for 60 min just after fertilization, which resulted in more than 30% of the hatched larvae being haploid androgenotes. They also observed the occurrence of a low number of diploid androgenotes. This new method provides new possibilities for artificial androgenesis without the need for irradiation in fish, thus simplifying and facilitating the induction procedure.
Although haploid androgenotes are inviable due to the expression of abnormalities, viability can be recovered by duplicating the haploid chromosome set using temperature or hydrostatic pressure shock at a time optimal for inhibiting the first mitotic division (cleavage division) [23]. By such induced endomitosis (chromosome doubling without cytokinesis), the resultant larvae are expected to become doubled haploids with complete homozygosity. However, such approaches to induce viable androgenetic doubled haploids have been considered to be technically very difficult [13]. Alternatively, several attempts have been made to induce diploid androgenotes using dispermy or diploid sperm in various fish species [24-28]. Nagoya et al. [29] successfully induced viable diploid androgenetic amago salmon (*Oncorhynchus masou ishikawae*) with gamma-irradiated eggs and subsequent dispermy fertilization, using sperm fused by polyethylene glycol. In rainbow trout (*O. mykiss*), Thorgaard et al. [30] reported improved survival of androgenetic diploids that were produced using sperm from tetraploid males. Recently, we successfully produced viable diploid androgenetic loach (*M. anguillicaudatus*) using UV-irradiated eggs and the cryopreserved diploid sperm of a neo-tetraploid male, which was produced by inhibition of the release of the second polar body after fertilization between a diploid female and a natural tetraploid male [21].

Here, we describe the procedure for artificial induction of diploid androgenotes by combining the cold-shock method and the use of diploid sperm in loach. Namely, eggs fertilized with diploid sperm from the neo-tetraploid male were immediately
cold-shocked to induce androgenesis just after fertilization. We identified the diploid status of the hatched larvae by flow cytometry, and also verified the all-male inheritance of the resultant diploid androgenotes by microsatellite DNA genotyping.

2. Materials and methods

2.1. Ethics

This study was performed according to the Guide for the Care and Use of Laboratory Animals in Hokkaido University.

2.2. Fish and gamete collection

Adult wild-type diploid female loaches were obtained from Iwamizawa city (Hokkaido, Japan). To induce neo-tetraploid males, the eggs of a normal diploid female were fertilized with diploid sperm from a natural tetraploid male, which were then heat-shocked (42 °C, 2 min) at 5 min after fertilization to inhibit second polar body release [31]. The experimental fish were reared in the aquarium room of the Environment Control Experiment Building of the Faculty of Fisheries Sciences, Hokkaido University. Eggs from a single female were used for cold-shock, UV-irradiation, and intact control treatments in each batch. A total of three females
and one neo-tetraploid male were used in the experiment.

Ovulation and spermiation were induced by the injection of human chorionic gonadotropin (20 IU/g body weight, Asuka Pharmaceutical Co. Ltd., Tokyo, Japan), as described by Suzuki and Yamaguchi [32]. After rearing at 27 (± 0.5) °C for 10-12 h, the fish were anesthetized in 0.1% 2-phenoxyethanol and the gametes were collected according to methods described in Morishima et al. [22].

2.3. Androgenesis by UV-irradiated eggs

Androgenesis induced by UV-irradiated eggs were used as controls in this study. Eggs were irradiated according to methods described in Fujimoto et al. [19]. For each batch, approximately 180 eggs were stripped onto 2 mL of seminal plasma prepared from the sperm of masu salmon (O. masou), which were then UV-irradiated at 150 mJ/cm².

Diploid sperm were added to the irradiated eggs, mixed well, activated by 20 (± 0.5) °C ambient tap water, and transferred to a container (the bottom of a 320 mL volume plastic box was cut and covered with mesh), which was placed in a Styrofoam box containing with 20 (± 0.5) °C ambient tap water.

2.4. Androgenesis by cold-shock treatment
Intact diploid sperm were added to the intact eggs, mixed well, and activated by 20 (± 0.5) °C ambient tap water. Just after activation (within 10 sec), eggs were quickly transferred to a container in a Styrofoam box containing 6 L of cold tap water at 3 (± 0.5) °C for 30 min. After the treatment, the container with the eggs was transferred to another box containing ambient tap water at 20 (± 0.5) °C.

2.5. Preparation of the control group

Eggs fertilized with intact diploid sperm without cold-shock treatment were used as the intact control group, and these eggs were transferred to a container in a Styrofoam box with ambient tap water at 20 (± 0.5) °C. At 150 min after fertilization, the eggs of the UV-irradiated group, the cold-shocked group, and the intact group were transferred to 90 mm plastic dishes, and incubated at 22 °C.

2.6. Frequencies of fertilized eggs, hatched larvae, and normal larvae

The fertilization rate was calculated as the frequency of cleaved eggs to total eggs used at 4 h after fertilization. The hatching rate was calculated as the proportion of hatched larvae to total eggs used. The normal rate was calculated as the proportion of normal hatched larvae to total eggs used.
2.7. Ploidy and paternity

Ploidy of larvae were analyzed at approximately 72 h after fertilization by flow cytometry (PA-II, Partec GmbH, Münster, Germany), according to the procedure described in Fujimoto et al. [19]. Each larva was first digested by 85 µL of solution A (CyStain DNA 2step, Partec GmbH, Germany) for 15 min, then 15 µL of the digested solution was mixed with 500 µL of solution B (CyStain DNA 2step, Partec GmbH, Germany), and analyzed by the flow cytometer. The remaining 70 µL of digested solution was used for DNA extraction for microsatellite genotyping. All hatched larvae from the cold-shocked and UV-irradiated groups were analyzed for ploidy status. In the intact control groups, ploidy status was determined for 50 normal and ten abnormal larvae.

Twenty-one diploid, ten triploid, and six tetraploid larvae appeared in the cold-shocked group, which were genetically analyzed, along with the female and male parents, using five loach microsatellite DNA loci: Mac 204, Mac 229, Mac 612, Mac 628, and Mac 638. The DNA extraction and microsatellite genotyping were performed following the methods of Morishima et al. [33].

2.8. Statistical analysis
The data are shown as mean ± S.D., based on triplicate experiments, and were analyzed with one-way ANOVA followed by Duncan’s multiple comparisons tests ($P > 0.05$) using R software.

3. Results

When intact eggs were fertilized with diploid sperm from the neo-tetraploid male and cold-shocked at $3 (± 0.5)$ °C for 30 min, the fertilization rate was $40.76 ± 3.41\%$, which was significantly lower than that of the intact control group ($P < 0.05$) (Table 1). In contrast, there was no significant difference between the fertilization rates of the cold-shocked group and the UV-irradiated group ($26.64 ± 18.85\%; P > 0.05$). The hatching rate of the cold-shocked group decreased abruptly to $29.63 ± 5.94\%$, which was 10% less than the fertilization rate; however, this type of decrease was not observed in the intact or UV-irradiated groups. Differences in hatching and normal rates among groups followed similar trends as the fertilization rate (Table 1).

Ploidy analysis confirmed the tetraploidy of the neo-tetraploid male (Figure 1A), and diploidy of the sperm spawned by that male (Figure 1B). The ploidy of all hatched normal and abnormal larvae from the cold-shocked and UV-irradiated groups were analyzed, along with 50 normal and ten abnormal larvae from the intact control group (Table 2 and Figure 1C-F). In the intact control group, besides the triploid, a few haploid, diploid and aneuploid larvae were detected. Although the majority of
hatched larvae from the UV-irradiated group were diploid, three triploid larvae and one mosaic abnormal larva also appeared. In the cold-shocked group, the larvae showed various ploidy states. Besides diploids, triploids, tetraploids, mosaics, and aneuploids, hexaploid and octaploid larvae were also detected in the cold-shocked group.

Diploid larvae were presumably putative androgenetic larvae, and the yield rate was calculated as the proportion of diploid larvae relative to total eggs used. For the cold-shocked and UV-irradiated groups, the yield rates of androgenetic diploids were $12.29 \pm 3.25\%$ and $22.23 \pm 13.42\%$, respectively, which were not significantly different ($P > 0.05$).

Paternity or all-male inheritance of the putative androgenetic diploid was analyzed with five independent microsatellite DNA loci. Diploid ($n = 21$), triploid ($n = 10$), and tetraploid ($n = 6$) larvae from one cold-shocked group were analyzed together with the female and male parent. Maternally and paternally derived alleles were detected in all triploid and tetraploid larvae. In contrast, only paternal alleles were detected in the diploid larvae (Table 3). For example, in the locus Mac229 from linkage group (LG) 3 [33], the female had the genotype 206/223, and the male had the genotype 176/190/219/260. Among the 21 diploid progeny, five progeny showed the 176/190 genotype, eleven progeny showed the 190/219 genotype, two progeny showed the 219/260 genotype, and the other three progeny had genotypes of 176/219, 176/260, and 190/260, respectively. The female-specific alleles, 206 and 223, appeared in the
triploid and tetraploid progeny.

4. Discussion

The use of diploid sperm improved the production of viable diploid androgenotes relative to that of androgenetic doubled haploids that were induced in genetically inactivated eggs fertilized with haploid sperm followed by induction of endomitosis at the first cleavage. In Misgurnus loach, the yield of doubled haploid androgenotes was less than 1%, and most could not survive beyond the early feeding stage [34]. In contrast, Arai et al. [24] reported higher survival rates (5.6 to 9.3%) in diploid androgenotes that were produced using diploid sperm of natural tetraploid and UV-irradiated eggs. Similar survival rates (7.14 ± 6.29%) were also reported when cryopreserved diploid sperm of neo-tetraploid males were used [21]. Higher survival rates of diploid androgenotes induced with diploid sperm from artificial tetraploids have also been demonstrated in rainbow trout [30]. However, the yield and survival rate of androgenotes are nonetheless lower relative to those of normal fertilization. Many variables can account for the low viability of androgenotes, such as egg quality, side effects of irradiation or cold-shock, and the heat-shock or pressure-shock administered at the prometaphase of the first mitosis. In addition, the expression of lethal genes due to the unmasking of homozygosity could also decrease the viability of androgenotes, especially in the case of doubled haploids.
In the present study, viable androgenetic diploid loaches were successfully induced by cold-shock treatment of eggs that were fertilized with diploid sperm. In general, natural or artificially induced tetraploid males generate diploid sperm. Some diploid-triploid mosaic [35] and masculinized clonal loach individuals also spawn diploid sperm [36-38]. These diploid sperm can be cryopreserved and used to restore viable diploid progeny through androgenesis, thus enabling the recovery of special fish strains or endangered species [21,30]. The androgenesis induction method we reported here may have potential applications in such restoring operations, and further offers a more convenient procedure in which to do so.

In summary, intact loach eggs that were fertilized with diploid sperm from neo-tetraploid male and then cold-shocked at 3 (± 0.5) °C for 30 min successfully produced viable androgenetic diploids. The exclusive paternal inheritance in these diploid larvae was genetically confirmed by microsatellite genotyping. These results indicate that cold-shock treatment just after fertilization induces androgenesis as effectively as does fertilization with UV-irradiated eggs.

Acknowledgements

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Misgurnus anguillicaudatus from UV irradiated eggs by suppression of the first


**Figure captions**

Figure 1. Relative DNA content. A: neo-4N; B: 2N sperm; C: 2N larvae of cold-shock group; D: 3N larvae of intact control group; E: 2.8N abnormal aneuploid larva of cold-shock group; F: 2.6N-3N abnormal mosaic larva of cold-shock group.
Table 1 Percent of fertilization, hatching, normal, and diploid larvae in the different treatment groups

<table>
<thead>
<tr>
<th>Female</th>
<th>Male</th>
<th>Treatment</th>
<th>No. of eggs</th>
<th>Fertilization (%)</th>
<th>Hatching (%)</th>
<th>Normal (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2N wild-type</td>
<td>Neo-4N</td>
<td>—</td>
<td>179 ± 7</td>
<td>63.98 ± 2.01</td>
<td>60.80 ± 3.85</td>
<td>57.87 ± 5.04</td>
</tr>
<tr>
<td>2N wild-type</td>
<td>Neo-4N</td>
<td>3 (± 0.5) °C, 30 min</td>
<td>182 ± 12</td>
<td>40.76 ± 3.41</td>
<td>29.63 ± 5.94</td>
<td>20.11 ± 1.16</td>
</tr>
<tr>
<td>UV eggs</td>
<td>Neo-4N</td>
<td>UV-irradiation</td>
<td>182 ± 24</td>
<td>26.64 ± 18.85</td>
<td>22.90 ± 16.74</td>
<td>18.46 ± 15.42</td>
</tr>
</tbody>
</table>

Different superscript letters within a column represent significantly different means (P < 0.05)
<table>
<thead>
<tr>
<th>Female</th>
<th>Male</th>
<th>Treatment</th>
<th>External appearance</th>
<th>Larvae no.</th>
<th>Ploidy status</th>
<th>Mosaic</th>
<th>Aneuploid</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>2N wild-type</td>
<td>Neo-4N</td>
<td>—</td>
<td>Normal</td>
<td>50</td>
<td>0 0 49 0 0</td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Abnormal</td>
<td>10</td>
<td>2 1 5 0 0</td>
<td>2</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>2N wild-type</td>
<td>Neo-4N</td>
<td>3 (± 0.5) °C, 30 min</td>
<td>Normal</td>
<td>110</td>
<td>0 51 36 19 3 1</td>
<td>3</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Abnormal</td>
<td>53</td>
<td>0 17 20 2 3</td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>UV eggs</td>
<td>Neo-4N</td>
<td>UV-irradiation</td>
<td>Normal</td>
<td>107</td>
<td>0 107 0 0 0</td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Abnormal</td>
<td>29</td>
<td>0 25 3 0 1</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

1: one 2.6N; 2: one 2.7N, one 2.8N; 3: one 2N-3N, one 2N-2.6N, one 2N-4N; 4: one 2.7N; 5: one 2N-2.4N, one 2.6N-3N, one 2.3N-3N; 6: one 2.3N, two 2.4N, one 2.6N, three 2.8N, one 3.5N, one 5.6N; 7: one 6N, one 8N; 8: one 2N-2.2N.
Table 3. Genotypes of diploid, triploid, and tetraploid progeny from the cold-shock treatment

<table>
<thead>
<tr>
<th>Locus (LG)*</th>
<th>Female</th>
<th>Male</th>
<th>Progeny from cold shock treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Diploid</td>
</tr>
<tr>
<td>Mac204 (LG9)</td>
<td>290/290</td>
<td>265/271/285/295</td>
<td>265/271: 5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>271/285: 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>271/295: 3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>285/295: 3</td>
</tr>
<tr>
<td>Mac229 (LG3)</td>
<td>206/223</td>
<td>176/190/219/260</td>
<td>176/190: 5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>190/219: 11</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>190/260: 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>219/260: 2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>160/180/213: 4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>230/230: 8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>165/165/187: 5</td>
</tr>
</tbody>
</table>

*: see Morishima et al. [33]