Artificial sperm insemination in externally fertilised fish as a novel tool for ex situ and in situ conservation of valuable populations

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ABSTRACT: Loss of genetic diversity and accumulation of deleterious mutations may lead to inbreeding depression in captive breeding. To address the problem of maintaining genetic diversity, we developed a new fish spawning method which offers flexibility in crossing diverse species when in vitro fertilisation (IVF) is not available. This method involves the collection of sperm from several males of ovuliparous fish; the sperm mix is then injected by catheter into the ovarian cavity of a female through the oviduct. We demonstrate, using zebrafish as a model for externally fertilised fish, that the sperm survives the ovarian conditions and can fertilise ovulated eggs, which are released from the body cavity during natural spawning. Wild type females were injected with reporter transgenic sperm from homozygous transgenic males before intended spawning with wild type males. The sperm injection method did not have an impact on reproductive parameters such as egg production or fertilisation rate compared to controls. In 25 successful spawning experiments, 20 females produced mixed genotype offspring comprising both transgenic and wild type larvae in varying ratios, indicating that the injected transgenic sperm efficiently competed with sperm released by non-transgenic wild type mating males, and both sperm types contributed to the fertilisation of the released eggs. This experiment provides proof of principle for increasing the genetic base of offspring of fish species, including that of many endangered fish species for which IVF is not available due to lack of timed induction of ovulation or when gametic release cannot be synchronised.

KEY WORDS: Genetic diversity · Sperm ovarian lavage · Zebrafish · Danio rerio

1. INTRODUCTION

Breeding fish in captivity (ex situ conservation) is a widely used management practice for endangered aquatic organisms. The main aim of such operations is to restock habitats and/or to restore vulnerable popu-
Therefore, alternative breeding methods are being made towards genetically selective breeding operations of fish intended for restocking to natural waters (Boscari & Congiu 2014), as well as developing protocols for specific training of the offspring to adapt to natural environments (Brown & Day 2002). However, such operations lack standardised procedures, and training fish in captivity remains problematic. Moreover, when schools of fish are held together, it is almost impossible to control the genotypes in the breeding process as there is no control over which fish will spawn within the enclosed group. Additionally, in such a case it is not possible to use cryopreserved sperm, which is an important tool in maintaining the genetic purity and diversity of the supported population. Another problem is that limitations on cage sizes influence the number of parents that can be used in paired- and group-spawning fish species. To address these concerns, a novel method of fish propagation is necessary in which the advantages of natural spawning efficiency can be combined with controlling genotype diversity. To achieve this goal, new technology for the control of genotypes is required, and gametes for the fertilisation of endangered species need to be made available of available gametes for fertilisation in endangered species needs to be overcome.

Munehara et al. (1989) described a subcategory of ovuliparous reproduction in elkhorn sculpin Alcichthys elongatus. The reproductive strategy of this species is internal gametic association (IGA), which refers to the association of male and female gametes inside the female reproductive tract followed by their release and subsequent fertilisation in the external environment. Müller et al. (2018a) imitated this strategy in an artificial breeding context, whereby collected sperm of ovuliparous common carp Cyprinus carpio were inserted into females by ovarian lavage and successfully contributed to fertilising released eggs. In that study, sperm samples were collected and injected into the ovary lobe with a catheter 12 h before ovulation in parallel with final hormonal induction. In another experiment (Müller et al. 2018b), a sperm and hormone mixture (carp pituitary extract, CPE) was injected into the ovary lobes of African catfish Clarias garepinus 10 h before ovulation. The absorbed CPE induced ovulation and all females produced good quality eggs that developed normally after water activation of the gametes and fertilization.

In neither of these experiments did ovarian fluid activate the spermatozoa, which thus maintained its biological activity for 10–12 h. These experiments demonstrate a proof of principle alternative to induced spawning. We hypothesise that genome diversity could be increased in pair-spawned or cohort-spawned fish by using ovarian lavage with mixed sperm samples. However, to prove this hypothesis, it is necessary to address the potential contribution to fertilisation of sperm stored in the ovary, on the one hand, and the native sperm from mating males, on the other.

In this study, our aim was to compare the fertilisation capability of naturally released and injected sperm in wild spawning fish. We chose zebrafish Danio rerio as a model species, as it is a well-established animal model for a wide range of research areas including reproduction biology and one in which mixing and monitoring of controlled genotypes are conveniently offered by transgenic reporter gene activity in offspring (Csenki et al. 2010).

2. MATERIALS AND METHODS

2.1. Zebrafish lines and husbandry conditions

Two zebrafish lines were used for the experiment. The wild-type zebrafish AB line has been bred for
several years in the Zebrafish Laboratory of the Hungarian University of Agriculture and Life Sciences. The transgenic zebrafish line Tg-2.4shh:gfpABC, carrying regulatory elements of the sonic hedgehog gene, was received from the Karlsruhe Institute of Technology (Ertzer et al. 2007) and has been reared under the same laboratory conditions as the AB line. Experimental AB fish were maintained in 1 l polycarbonate tanks at a ratio of 1 female to 2 males. Transgenic males were kept together (n = 15, standard length [SL]: 29 ± 1.6 mm) in a 3 l polycarbonate tank with a water flow recirculated system (ZebTec, Tecniplast) through an upwelling bead filter at 25 ± 2°C and were fed 3 times per day with commercial flakes (Sparos Zebra; 400−600 μm) and live Artemia larvae grown from cysts (Ocean Nutrition; >230 000 nauplii g−1). The photoperiod was set at 14 h light:10 h dark.

The protocols of fish propagation and the template informed consent forms (Scientific Ethics Council for Animal Experimentation; XIV-001-2306-4/2012 and PE/EA/742-7/2020) were reviewed and approved by the Hungarian National Food Chain Safety Office, Animal Health and Animal Welfare Directorate of the Government Office of Pest County with respect to scientific content and compliance with applicable research subject regulations.

2.2. Spawning

To confirm the identification of offspring genotype, 2 × 15 AB females (SL = 26.1 ± 1.3 mm) and 2 × 30 AB males (SL = 26.6 ± 1.3 mm) were selected randomly. The AB line began from unknown zebrafish source stocks bought from 2 pet shops (pet shop A and pet shop B) in Albany, Oregon, USA, in the early 1970s. Haploid progeny from AB females were crossed with random AB males for approximately 70 generations until the early 1990s, when 6 diploid progeny stocks (each from a distinct haploid female) were thoroughly intercrossed to produce the modern AB line. The current AB source stock is maintained through large group-spawning crosses (Holden & Brown 2018).

Traditional spawning approaches were used. Specifically, zebrafish propagation took place in ~1.7 l spawning tanks (Sloping Breeding Tank; ZebTec, Tecniplast) which feature a sloped interior or ‘beach style’ that facilitates and promotes zebrafish spawning (these tanks were used for all treatments). The tanks contain 2 interchangeable containers; the bottom of the inner vessel is perforated to facilitate egg collection and movement of fish post-spawning. A schematic of the breeding design is shown in Fig. 1. In the 1st spawning treatment (Spawning I), 1 female and 1 male were introduced into each breeding tank; in Spawnings II, III and IV, 1 female and 2 males were placed into each breeding tank. Spawning took place the following morning, typically a few hours after the lights were turned on, as zebrafish reproduction is strongly influenced by photoperiod (Nasiadka & Clark 2012). All fish were released to spawn according to the protocol of zebrafish fertilization and embryo isolation (www.zfic.org/common%20techniques/mating.pdf). For all experiments, 1 l breeding tanks (ZebTec, Tecniplast) were used. Water conditions were as follows: temperature 25°C; pH 7.0 ± 0.2; average conductivity 525 μS. Eggs were collected from every tank and placed into Petri dishes (diameter: 100 mm) 2 h after the start of the light photoperiod. Eggs were incubated in a thermostat (25.5°C, photoperiod was set at 14 h light:10 h dark) with daily water changes. After a 72 h incubation period, all eggs were checked and imaged with a Leica M205 FA microscope.

2.3. Artificial insemination

Tg-2.4shh:gfpABC homozygous males were removed from the breeding tank and anaesthetised with MS-222 (4.2 ml MS-222 per 100 ml system water). After anaesthetisation, the genital area was dried with a paper towel and sperm stripping occurred under a Leica M205 FA microscope. Sperm was obtained by gentle pressure on the sides of the Tg-2.4shh:gfpABC males and collected using pipette tip (20−200 μl, Gilson, thermo, yellow) and a half-length of G-1 glass capillary (length 90 mm, external diameter 1 mm, internal diameter 0.6 mm; Nari-shige Scientific Instrument). One male’s sperm sample (~1 μl, 0.4−1.4 μl) was artificially inseminated into 1 AB female. Females were anaesthetised in the same way as males 1 h from the expected spawning time (dark/light changes). A glass capillary was inserted approximately 2 mm deep into the oviduct through the genital papilla of anaesthetised females using an automated pipette. Sperm samples were injected into the centre of the genital papille, i.e. sperm distribution was not directed into the 2 oviducts (random distribution), and then females were put back into the spawning tanks with AB males (Fig. 1). Resting periods varied among the spawning times depending on the experimental conditions and possibilities (Table 1).
2.4. Analysis of offspring genotype

The genotypes of the freshly hatched larvae were investigated using a Leica M205 FA microscope with Leica Application Suite X v.3.4.2.18368 software (LAS X; Leica Microsystems). Offspring originating from transgenic sperm were counted upon detection of green fluorescence protein (GFP) expression using a GFP2 long pass filter, where the maximal excitation and emission values were 489 and 508 nm, respectively. Females that either did not provide eggs/larvae from sperm insemination by the 3rd and 4th experimental cycles (females 8, 12, 14, 29) or died during the experiments (females 4, 10, 20, 24) were removed from further analysis.

Table 1. Intervals of the zebrafish experimental spawning series (in days) (see Section 2.2 and Fig. 1)

| No. of females | Spawning | I | II | III | IV |
|----------------|----------|---|----|-----|----|
| 1−15           |          | 0 | 14 | 7   | 14 |
| 16−30          |          | 0 | 7  | 15  | 25 |

2.5. Statistical analysis

Statistical analyses of fertilisation rates and egg number female\(^{-1}\) values were carried out by 1-way
ANOVA (with Tukey post hoc test) in SPSS v.22 for Windows. Treatment means were compared using a significance level of $\alpha = 0.05$. The correlation between the ratio of fertilised egg yield to hatched transgenic larva was analysed by logarithmic correlation to determine the line of best fit to the data.

### 3. RESULTS

Our first question was whether the zebrafish sperm insemination protocol influenced the number of eggs released from treated females. As shown in Table 2, there was no statistical difference in the number of eggs released between the controls and Spawning III treatment individuals. Thus, sperm insemination did not influence the spawning success of the same female spawning naturally or after sperm injection. In the Spawning IV treatment, the females that were inseminated with sperm released more eggs than on previous occasions, but there was no significant difference in case of female was introduced with 1 or 2 males in spawning tanks (Table 2). There were no statistical differences ($p < 0.05$) among the total fertilisation rates independent of spawning method (natural spawning – artificial insemination) when the female was introduced with 1 or with 2 males in the spawning tanks (see Table 2).

Next, we investigated if sperm from naturally mating males successfully fertilised eggs during wild spawning or if the eggs were fertilised by artificially inseminated transgenic sperm. To answer this question, we examined the phenotypic appearance of transgenic larvae, which are identified by GFP fluorescence by the emerging cells of larvae in which the sonic hedgehog regulatory elements activate the fluorescent reporter gene. Prior to the experiments, we confirmed that the transgenic males were homozygous and that all of their offspring inherited a copy of the expressed transgene (Fig. 2 and data not shown). We then confirmed that the control wild type males were indeed void of any fluorescence signal at the protruding mouth stage (72 h post-fertilisation [hpf]) (Fig. 2). Finally, we carried out spawning experiments with the injected females and analysed their offspring for transgene activity at the protruding mouth stage (Table 2, Fig. 2).

As demonstrated in Table 3, there were 25 successful spawnings that resulted in larvae and 20 females that produced offspring including both transgenic-reporter-positive larvae and AB wild type larvae in varying fertilisation ratios. The average percentages of hatched transgenetic larva were $31.3 \pm 29.1\%$ (all treated females: $n = 25$, min.–max.: $0–84.3\%$) and $39.1 \pm 27.4\%$ (all treated females which produced living embryos: $n = 20$, min.–max.: $3.8–84.3\%$). In 5 spawnings (20%) there were no transgenic larvae produced from the eggs.

### 4. DISCUSSION

In this study, we provide evidence for the generation of offspring with a mixed genetic background, including both transgenic and non-transgenic larvae originating from artificial insemination of transgenic fish sperm into wild type females followed by natural crossing with wild type, non-transgenic males. This experiment, carried out with zebrafish, provides proof of principle evidence for how genetic diversity of offspring could be increased by using mixed sperm insemination in an externally fertilising fish species, and we propose that this approach could be applied in similar externally fertilising fish species that are identified for genetic conservation efforts. Additionally, breeding programs aiming to increase the genetic diversity of economically important fish species, which are propagated through induced

| Treatment | Spawning experiment (no. of spawned females) | Control | Spawning II + Spawning III control (n = 14) | Spawning III (n = 15) | Spawning IV (n = 13) |
|-----------|---------------------------------------------|---------|--------------------------------------------|----------------------|----------------------|
| Sex ratio | $1 \varphi \times 1 \sigma'$ | $1 \varphi \times 2 \sigma'$ | $1 \varphi \times 2 \sigma'$ | $1 \varphi \times 2 \sigma'$ |
| No. of eggs released (mean ± SD) | $34.8 \pm 27.3^a$ | $68.6 \pm 36.7^{ab}$ | $62.5 \pm 39.7^{ab}$ | $97 \pm 81.8^{b}$ |
| Fertilisation rate (%) (mean ± SD) | $21.7 \pm 30.7^a$ | $43.3 \pm 29.4^{b}$ | $42.9 \pm 27.5^{b}$ | $41.5 \pm 23.7^{b}$ |
spawning could also benefit from our approach by using selected high-quality cryopreserved sperm samples and allowing their delivery to females which require pairing or group spawning to induce ovulation.

Our results show that the inseminated transgenic sperm takes part in the fertilisation process together with the sperm of the naturally spawning male and results in a mixture of both transgenic and non-transgenic larvae, with a proportion ranging from 0–84.3% transgenics. The reason for this large range may lie in suboptimal technical factors as well as yet unidentified differences in sperm quality or fitness resulting in sperm competition (Taborsky 1998). Such parameters may also include unidentified technical errors in methodology, such as inadvertent water contamination of sperm during handling, leading to premature activation of spermatozoa before ovulation of target eggs. Alternatively, suboptimal inseminated sperm delivery into the oviduct could lead to sperm leakage/release from the inseminated females. Based on preliminary observations, there are several technological parameters that could be improved before adaptation to other fish species.

Injected sperm does not cause internal fertilization. In ovuliparous fish species, eggs are released from the female’s body followed by their subsequent external fertilization or activation in the water. Dean et al. (2019) were the first to find living embryos in a non-copulatory, egg-laying teleost species (three-spined stickleback Gasterosteus aculeatus) and remove them from the ovary lobes at the eyed stage. Larvae were hatched and reared normally to adulthood under controlled conditions. Munehara et al. (1989) described a subcategory of ovuliparity, called IGA, which refers to the association of male and female gametes in the female reproductive tract followed by their release and subsequent fertilization in the external environment. Müller et al. (2020) investigated the sperm–ova interaction just after gamete stripping without water activation in the African catfish Clarias gariepinus using an electron microscope. Spermatozoa were distributed near the micropylar region and detected within the micropyle canal, similar to the observations of Munehara et al. (1989). We hypothesise that the spermatozoa are inactive near the micropyle region (or in it), and the closest sperm cell to the micropyle just after gamete release will fertilise the egg after water activation outside the female’s body.
In this study, freshly stripped sperm were injected into the ovary a few hours before the expected spawning time. We previously demonstrated that sperm insemination can be applied successfully 12 h before ovulation in common carp *Cyprinus carpio* and 10−36 h before ovulation in African catfish *C. gariepinus* (Müller et al. 2018a,b, 2019, 2020). In these studies, the potential effects of different latency times of the inseminated spermatozoa were investigated. However, there was no statistical difference in fertilisation and hatching rates between 5 and 36 h of latency in African catfish (Müller et al. 2020), indicating a surprisingly elongated viability and/or fertilisation capacity of the inseminated spermatozoa. The latency times used here are expected to be applicable to other fish species as well and offer a suitable time window for programmed spawning upon various hormonal treatments. Cryopreserved sperm can be used in this method (Müller et al. 2019).

To illustrate the potential advantages of the insemination approach in traditional fish rescue programs (*in situ* conservation) with other species, a schematic representation of the genotype combinations and comparisons of reproduction parameters are shown in Fig. 3.

| No. of Spawning III | AB ♀ (Tg sperm inj.) × 2 AB ♂ | 72 hpf (%) | Tg larvae (%) |
|---------------------|---------------------------------|------------|---------------|
| 1 157               | 54.1                            | 48.2       |
| 2 31                | 83.9                            | 3.8        |
| 3 –                 | –                               | –          |
| 5 15                | 60.0                            | 0          |
| 6 –                 | –                               | –          |
| 7 –                 | –                               | –          |
| 9 30                | 60.0                            | 16.7       |
| 11 99               | 2.0                             | 0          |
| 13 27               | 14.8                            | 25.0       |
| 15 28               | 25.0                            | 14.3       |
| 18 93               | 64.5                            | 46.7       |
| 19 61               | 72.1                            | 9.1        |
| 21 –                | –                               | –          |
| 22 121              | 33.9                            | 53.7       |
| 23 –                | –                               | –          |
| 25 60               | 67.0                            | 75.0       |
| 26 65               | 66.2                            | 0          |
| 27 61               | 29.5                            | 11.1       |
| 30 48               | 2.1                             | 0          |

| No. of Spawning IV | AB ♀ (Tg sperm inj.) × 2 AB ♂ | 72 hpf (%) | Tg larvae (%) |
|--------------------|---------------------------------|------------|---------------|
| 1 143              | 87.4                            | 24.8       |
| 2 71               | 54.9                            | 20.5       |
| 3 48               | 33.3                            | 81.3       |
| 5 63               | 33.3                            | 71.4       |
| 7 67               | 62.7                            | 28.6       |
| 9 60               | 16.7                            | 28.6       |
| 11 144             | 36.8                            | 9.4        |
| 13 71              | 46.7                            | 84.3       |
| 15 61              | 42.6                            | 77.1       |
| 16 144             | 11.1                            | 56.2       |
| 18 331             | 46.2                            | 84.3       |
| 19 61              | 42.6                            | 77.1       |
| 20 144             | 11.1                            | 56.2       |
| 21 331             | 46.2                            | 84.3       |
| 22 61              | 42.6                            | 77.1       |
| 23 61              | 42.6                            | 77.1       |
| 24 61              | 42.6                            | 77.1       |
| 25 61              | 42.6                            | 77.1       |
| 26 61              | 42.6                            | 77.1       |
| 27 61              | 42.6                            | 77.1       |
| 28 61              | 42.6                            | 77.1       |
the safety of larvae (Müller et al. 2019) is improved in the surrogate habitat (Tóth et al. 2016).

The disadvantage of the controlled spawning strategy is that the resulting genetic variability remains limited, as the genetic pool of offspring is limited to the availability and fecundity of parents selected from the original habitat (Fig. 3: crossing strategy × traditional fish rescue, controlled spawning program). Since males continue to produce sperm after stripping, they can be reintroduced back into their original habitat, where they can breed naturally (Fig. 3: crossing strategy × sperm insemination). By using the sperm injection method, expansion of the genetic diversity of the offspring can be achieved in a relatively small spawning area compared to a traditional fish rescue setup in which the number of parents used remains limited. Theoretically, an additional potential advantage of the sperm injection method is that, because of the lack of direct contact between males and females, no ectoparasite exchange between the 2 sexes originating from different populations will occur (Fig. 3: parasite transfer × traditional fish rescue, controlled spawning program). In contrast, conventional breeding and directional breeding strategies will increase the chance of parasite transfer, and thus these procedures may cause more harm than benefit. Some extreme examples of parasite transfer occurred between differently sensitised subspecies European carp and koi carp to Thelohanellus nikolskii (Myxosporea) infection (Molnár 2002) or Anguillicoloides crassus infection from Anguilla japonica to A. anguilla and A. rostrata (Sprengel & Lüchtenberg 1991, El-Shehabi et al. 2018).

4.1. Gene changes between neighbouring inbred populations

Endangered species often live in reduced or fragmented habitats, and as a result population fragmentation may occur, leading to inbreeding depression and a decrease in the genetic variability of the species. It is possible to increase the genetic diversity of isolated, inbred populations by applying the sperm injection method. In this procedure, males with excellent reproduction fitness are selected just before the spawning season and stripped of sperm; the sperm from several males is pooled and injected into...
one side of the ovarian lobe of anesthetised females. It is expected that when the females are released back to their home environment for natural spawning, eggs from the second lobe will be fertilised by males originating from the native habitat, while the injected lobe will release eggs that will be fertilised by the injected sperm. Thus, the genetic diversity of the native population is increased. There is no need for specialised equipment for short-term sperm storage because sperm quality parameters can be maintained within the first 10 h in cooling boxes (Pires et al. 2019). Only water contamination needs to be avoided. It is also possible to use sperm that has previously been cryopreserved and thawed for insemination, as was shown with *C. gariepinus* (Müller et al. 2019). The injected females can then be released where they were originally caught so that they can find their favoured spawning environment.

4.2. *Ex situ* conservation

The success of restocking programs for endangered species depends on the number and genetic diversity of the stocked individuals intended to maintain appropriate genetic variability (Ortega-Villaizan et al. 2011). *In vitro* fertilisation (IVF) and restocking numerous larvae or fingerlings could be an appropriate method to recover new or degraded habitats. IVF is generally used for *ex situ* conservation biology (captive breeding and rearing). Several fish species would benefit from such rescue efforts; however, IVF is not available for all species because fish may not respond to conventional hormonal injection, delicate broodstock may be sensitive to invasive hormone induction (Watson et al. 2009), or the fish may be too small (i.e. Vulnerable *Umbra krameri*, or Critically Endangered *Romanichthys valsanicola*, etc.). In some fish species like *U. krameri* there is no *in vitro* fertilisation technology available for propagation, and therefore the only possible method of artificial reproduction is induced spawning in a pen or tank (Kucska et al. 2016, Tatár et al. 2017). However, pen spawning can lead to bottleneck effects, as it strongly reduces the genetic diversity of the offspring due to the limited availability of mating pairs. The sperm insemination method, which allows a broad range of male genotypes to be used in fertilisation, may reduce the chance of such genetic deterioration.

In schooling fishes such as *Anguilla* spp., *in situ* conservation work is currently impossible due to the unique adaptation of the fish to inaccessible spawning environments. To circumvent this problem, several strategies are being developed to induce spawning of eel species in captivity. There are 2 types of induced propagation methods that are now available for European and Japanese eels in captivity. One is based on *in vitro* fertilisation and the other on induced spawning, whereby upon final hormonal administration females are placed into spawning tanks with 1–3 males to promote natural spawning. Horie et al. (2008) and Di Biase et al. (2016) reported that fertilisation and hatching were significantly higher with spontaneous spawning than with IVF. However, the success of induced spawning depends on brood preparation, which differs for the 2 sexes and may lead to variability due to differences in the quality of the sperm of successful male breeders. Spawning success is independent of the activation and fertilising ability of spawning males; thus, fertilisation success may be enhanced by combining previously selected, high-quality sperm for insemination of females which are then mated with well-conditioned males to enhance spawning success. The inseminated sperm is expected to be accumulated and to adhere to the micropyles (Müller et al. 2020), thus reducing the chance of spawning males outcompeting the inseminated sperm and thereby reducing the role of the spawning males in inducing ovulation. Additionally, the genetic basis of the offspring may be broadened by using pooled, preselected sperm samples from several males.

In this study, we have shown proof of principle for increasing the genetic diversity of offspring after sperm insemination in a genetic model species. We propose that our strategy should be tested on endangered species to prove our hypothesis in practice and to study how competition between sperm samples with different origins can contribute to the fertilisation of eggs released from the recipient females.

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