No support for cryptic choice by ovarian fluid in an external fertilizer

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Abstract
Whether the ovarian fluid (OF) represents a selective environment influencing cryptic female choice was tested using an external fertilizer experiencing intense sperm competition and large effects of OF on sperm swimming behavior—the Arctic charr (Salvelinus alpinus). We physically separated the OF from the eggs of reproductively active females and reintroduced either their own OF or fluid from another female to the eggs. The eggs were then fertilized in vitro in a replicated split-brood design with sperm from two males under synchronized sperm competition trials, while also measuring sperm velocity of the individual males in the individual OFs. We found large effects of males, but no effect of females (i.e., eggs) on paternity, determined from microsatellites. More important, we found no effect of OF treatments on the relative paternity of the two competing males in each pair. This experimental setup does not provide support for the hypothesis that OF plays an important role as medium for cryptic female choice in charr. Power analyses revealed that our sample size is large enough to detect medium-sized changes in relative paternity (medium-sized effect sizes), but not large enough to detect small changes in relative paternity. More studies are needed before a conclusion can be drawn about OF's potential influence on paternity under sperm competition—even in charr.

KEYWORDS
cryptic female choice, mate choice, ovarian fluid, Salvelinus alpinus, sperm competition

1 | INTRODUCTION

Sperm competition occurs when ejaculates from different males compete over fertilizing eggs either inside the females' reproductive tract or externally to the female's body (Birkhead & Møller, 1998; Parker, 1970; Pizzari & Parker, 1998; Simmons, 2001; Stockley, Gage, Parker, & Møller, 1997). For external fertilizers, the outcome of this competition, that is, which male successfully sire the offsprings, may be influenced by the relative competing abilities of the males' ejaculates (sperm and seminal fluids), the female's spawning products (eggs and ovarian fluid [OF]), and their potential interactions. Thus, females of external fertilizers might not be regarded as only providing an arena for sperm competition, as they may also actively discriminate among which of the males involved in sperm competition is allowed to fertilize her eggs through cryptic choice (Birkhead, 1998; Eberhard, 1996; Olsson, Shine, Madsen, Gullberg, & Tegelstrom, 1996; Thornhill, 1983; Zeh & Zeh, 1996). Mechanisms used by females to control which male sires her eggs are expected to evolve in species with multiple matings—particularly in species that are exposed to intense sperm competition where female control of which
males participate in the sperm competition is limited. This is the case for highly polyandrous broadcast spawners (Firman, Gasparini, Manier, & Pizzari, 2017), as for example in species with lek-like mating systems. In some such species, cryptic female control under sperm competition has the potential to generate large differences in offspring survival (Rudolfsen, Figenschou, Folstad, Nordeide, & Søreng, 2005; Wedekind, Muller, & Spicher, 2001).

In several of the first studies of external fertilizers documenting that sperm velocity is of importance for fertilization success under sperm competition, sperm velocity was measured in water (e.g., Gage et al., 2004; Liljedal, Rudolfsen, & Folstad, 2008; Rudolfsen, Figenschou, Folstad, Tveiten, & Figenschou, 2006; Ottesen, Babiak, & Dahle, 2009; Skjæraasen et al., 2009). However, eggs are embedded in OF, which is a semiviscous liquid, sometimes comprising 10%-30% of the volume spawned (Lahnsteiner, 2002). The OF has several important effects on sperm traits including extending sperm longevity and increasing sperm velocity (Butts, Johnson, Wilson, & Pitcher, 2012; Golpour, Esfandyari, & Dadras, 2012; Lahnsteiner, 2002; Litvak & Trippel, 1998; Turner & Montgomerie, 2002; Urbach, Folstad, & Rudolfsen, 2005). More important, there is, at least in some species, an interaction effect of OF on sperm velocity (Rosengrave, Gemmell, Metcalf, McBride, & Montgomerie, 2008; Urbach et al., 2005). That is, OF of individual females specifically promote sperm velocity of certain males over others, suggesting that OF may be a medium in which females exert cryptic female choice (Beirão, Purchase, Wringe, & Fleming, 2015; Dietrich et al., 2008; Nordeide, 2007; Rosengrave et al., 2008; Urbach et al., 2005). Recent studies on external fertilizers have demonstrated that OF may reduce the velocity of sperm from subordinate males relative to that of dominant males (Egeland, Rudolfsen, Nordeide, & Folstad, 2016; Lehnhert, Butts, et al., 2017; Makiguchi, Torao, Kojima, & Pitcher, 2016). Thus, subordinates, who invest more resources in their sperm and usually show the highest sperm velocity in water, have lower gains from their investments in sperm velocity than dominant males when sperm enter the OF surrounding eggs. Thus, females of external fertilizers may be promoting fertilizations by sperm from dominant males, not only by releasing their gonadal products close to and in synchrony with them, but also by promoting their sperm swimming performance in the immediate vicinity of the eggs.

Recent reviews and meta-analysis have, however, provided weak evidence for genetic benefits from polyandry (Slater, Mautz, Backwell, & Jennions, 2012) and few clear demonstrations of cryptic female choice (Firman et al., 2017). Whereas some studies on external fertilizers have reported increased fertilization success for the male whose sperm swims faster in the females’ OF (e.g. Evans, Rosengrave, Gasparini, & Gemmell, 2013; Rosengrave, Montgomery, & Gemmell, 2016; Lehnhert, Butts, et al., 2017; Lehnhert, Helou, Pitcher, Heath, & Heath, 2018), another study found no such relationship (Lehnhert, Heath, Devlin, & Pitcher, 2017). Additionally, no effect of cryptic female choice on offspring fitness was documented in salmon (Salmo salar) reported by Lumley et al. (2016). Our knowledge about known proximate mechanisms enabling females to affect the outcome of sperm competition is limited (Firman et al., 2017). Important exceptions are the role of the sperm protein binding in egg–sperm recognition in sea urchins (Palumbi, 1999), and egg glycoproteins’ role in avoiding inbreeding in mice (Firman & Simmons, 2015). The role of MHC-dependent gamete recognition might also be important in nonrandom gamete recognition and fusion in mice and salmonids (reviewed by Firman et al., 2017, Box 2). In Chinook salmon (Oncorhynchus tshawytscha), sperm from dominant males differ from those of subordinate males in flagellar beat frequency, bend length, bend angle, and wave amplitude when swimming in the OF (Butts, Prokopchuk, Kaspar, Cosson, & Pitcher, 2017). The fluid contains 174 proteins with individual variation in numbers and concentrations (Johnson et al., 2014) that may interact with the spermatozoa and modify their flagellar beating and velocity (Johnson et al., 2014; Rosengrave et al., 2016). Yet, although polyandrous and promiscuous females may derive fitness benefits from cryptic female choice, the general mechanisms are still unclear in the majority of taxa and species.

In external fertilizers like Salmonidae, it is not established whether the OF or the eggs themselves may influence the paternity during reproduction. Yet, the relative importance of these two factors might, at least in theory, nicely be identified by physically separating the OF from eggs of females and adding the OF from another female. These reproductive products could then be exposed to sperm competition trials, and the relative paternity from such trials could be compared to paternity when OF has not been exchanged between eggs from different females. Yeates et al. (2013) exchanged OF between eggs from salmon (S. salar) and brown trout (S. trutta) and reported that conspecific sperm gained fertilization precedence in interspecific sperm competition trials. Moreover, this precedence was primarily controlled by OF by increasing motility of conspecific sperm (Yeates et al., 2013). Yet, an intraspecific study on Chinook salmon (O. tshawytscha) documented no overall effect of OF on paternity success and no evidence for male–female interactions on paternity (Evans et al., 2013). This is surprising, as positive associations between sperm velocity in OF and both fertilization success and embryo survival have been reported in the same species (Rosengrave et al., 2016). Thus, the current experimental evidence suggests no intraspecific effect of OF on the outcome of sperm competition when exchanging OF between eggs from different females and exposing them to sperm competition.
away) when trying to guard the females from “sneakers” before spawning (Sigurjonsdottir & Gunnarsson, 1989). Yet, the spawning area provides no physical protection for the spawning pair and several males typically spawn in competition when the female releases her eggs (Sigurjonsdottir & Gunnarsson, 1989; Særum et al., 2011). In our studied charr population, 76.5% of the ejaculates experience sperm competition and the mean number of males releasing melt in each spawning event is 2.6, suggesting a high level of sperm competition (Særum et al., 2011). Sperm velocity and sperm density differ predictably between males adopting dominant or subordinate spawning strategies; that is, subordinates have more sperm with higher velocity in water, yet lower velocity in OF, than dominants (Egeland et al., 2016; Rudolfsen et al., 2006). Additionally, velocity of sperm in ejaculates has also been shown to influence fertilization success under sperm competition (Egeland, Rudolfsen, & JT, Folstad I., 2015; Liljedal et al., 2008).

In our present experiment, we conducted in vitro fertilization trials using charr gametes to disentangle the potential effects of eggs and OF in influencing relative paternity of two males under sperm competition. We first physically separated eggs and OF before embedding the eggs in either own or foreign OF. Thereafter, the eggs were fertilized by simultaneously releasing ejaculates from two males in competition, while also recording sperm speed in OF. If OF acts as a medium for cryptic female choice, we predicted that our experimental exchange of OF between eggs from the two females would influence paternity.

2 | MATERIALS AND METHODS

2.1 | Sampling and stripping of gametes

The charr were caught in Lake Fjellfrøsvatn located at 69°N, 19°E, at an altitude of 126 m in northern Norway, from the 18 to the 23 September in both years. Gillnets of 24 mm mesh size were used for fishing at three different spawning grounds (see Figenschou et al., 2004). Fish were removed from nets as soon as they were trapped to avoid injuries and thereafter stored in chicken wire cages by the shore until further handling. Males were caught less than 24 hr before they were stripped for gametes, whereas females—being more rare at the spawning grounds and hence more difficult to catch—were caught from 0 to 4 days prior to handling of the gametes. In the laboratory, the fish were put to death by a stroke to the head and fin tissue samples were obtained and kept in 70% ethanol for later genotyping. The area around the genital pore was then dried carefully by paper tissue in order to avoid contamination and subsequent activation of gametes, before the fish were stripped for free-running gametes by a gentle bilateral pressure from the anterior part of the abdomen toward the genital pore. Handling of the gametes and fertilizations were carried out by the same experienced individuals on the 23 and 24 September in both years.

2.2 | Experimental design

Two experiments were carried out in this study using a North Carolina II design. The first experiment (Sperm velocity analysis) was carried out to test whether the velocity of sperm from each of two males differed when swimming in each of two OF–water solutions from two different females. The details from this experiment are presented below (see Handling of male gametes). The second experiment (Paternity analysis) was performed to test for differences in relative paternity when sperm from two different males competed to fertilize eggs from a particular female which were surrounded by either its own OF or OF from another female. This second experiment was carried out in a block design where each block consisted of two males and two females that were tested in replicates (see Table 1). Females and males were randomly assigned to the blocks.

In each trial, eggs of one female were treated either as “control,” “own OF,” or “foreign OF.” For each treatment, sperm of both males

| TABLE 1 | The experimental design used to test effects of ovarian fluid (OF) and sperm identity on fertilization success. For each block, two males and two females were used. In each trial, sperm from two males competed to fertilize eggs of one female which were treated as either “control” (untreated eggs), “own OF” (own ovarian fluid removed and added again, i.e., treatment control), or “foreign OF” (ovarian fluid removed and replaced with that of the other female). Each treatment combination per trial was replicated for a total of eight blocks. Due to eggs of poor quality, we had to exclude one of the two trials in four of the eight blocks. Thus, our experiment consists of 72 observations (in vitro fertilizations), with 36 unique *male ID–egg donor ID–ovarian fluid treatments* (observations or fertilizations). The offspring of each treatment combination was genotyped to assess paternity and counted in order to determine the relative paternity for each trial and treatment. The total number of offspring genotyped was 649, whereas the mean number of genotyped offspring in each of the 72 fertilizations was 9.0. Since the fertilizations were carried out in two replicates (Table 1), the mean total number of genotyped offspring per pair of parents per treatment (control, own, and foreign) was 18.0.

| Predictor | Block | Trial 1 | Trial 2 |
|-----------|-------|---------|---------|
| Male      | Sperm ♂1 + ♂2 | Sperm ♂1 + ♂2 |
| Treatment | Control | Own OF | Foreign OF | Control | Own OF | Foreign OF |
| Egg-donor ID | eggs ♀1 + eggs ♀1 | eggs ♀1 + eggs ♀2 | eggs ♀2 + eggs ♀2 |
| OF ID     | OF ♀1 | OF ♀1 | OF ♀2 | OF ♀2 | OF ♀2 | OF ♀1 |
| Replicates | 2 | 2 | 2 | 2 | 2 | 2 |
competed to fertilize the eggs in two replicates. We started out the experiment with eight full blocks, each including two trials as in Table 1, which resulted in 16 trials and the total sample size of 96 observations (in vitro fertilizations with 48 unique combinations of male-egg donor-OF treatments, see below). However, in four of the blocks, one of the two females (i.e., one of the two trials in the block) produced eggs of poor quality and no eggs survived in one or more of the replicates. We therefore had to exclude one of the trials in each of these four blocks. Thus, our experiment consisted of four full blocks, each with 12 observations, and four half blocks, each with six observations. This gives a total of 72 observations (in vitro fertilizations with 36 unique combinations of male-egg donor-OF treatments, see below). A total of 649 offspring were genotyped, whereas the mean number of genotyped offspring in each the 72 fertilizations was 9.0. Since the fertilizations were carried out in two replicates (Table 1), the average total number of genotyped offspring per pair of parents per treatment (control, own and foreign) was 18.0 (minimum and maximum are 9 and 20 offspring, respectively).

Fertilization success was measured as the ratio of offspring sired by the focal male (assigned to the male with the lower id number) to the sum of offspring of both males (termed “relative paternity”). "Relative sperm velocity" (i.e., "VCLdiff") was measured as the difference in sperm velocity between the focal male and the competing male.

2.3 Handling of male gametes

All handling of gametes was conducted in a precooled laboratory. Immediately after stripping the milt from each male in a petri dish, we estimated milt volume (in 1-ml syringes to the nearest 0.1 ml) and spermatocrit. When not handled, the milt was kept at 4°C in closed 1.5-ml Eppendorf tubes and potential effects of handling time of sperm were minimized by conducting all the ejaculate measurements in the order by which the fish were included in the experiment. Sperm behavior was recorded 10 s postactivation, that is, as fast as possible. Sperm behavior was recorded first in lake-water to ensure that the sperm were active, and then in water-diluted OF (ratio of OF to water was 1 to 2, or 33% OF) of the two females in each block (see below). The same or similar dilution of water and OF was used in previous studies (e.g., Butts et al. 2012; Egeland et al., 2015; Egeland et al., 2016; Urbach et al., 2005). For simplicity, we hereafter refer to this mixture as OF. After placing less than 0.12 µl of sperm on a precooled (5–6°C) standard counting chamber (Leja Products BV), measurements were initiated after activating sperm by adding 4.5 µl of water or OF. To avoid possible effects of cell density on sperm behaviors, the slides were, immediately after sperm activation, screened for areas with an appropriate density of cells (average number of motile sperm was 97, SD = 42.4, N = 32). Records were made using a CCD black and white video camera module (Sony, XC-ST50CE) attached to a CH30 Olympus microscope with a negative phase-control objective lens (×10 magnification). All recordings were carried out in replicates. The video recordings were analyzed using an HTM-CEROS sperm tracker (CEROS version 12; Hamilton Thorn Research, Beverly, MA), a standardized computer-assisted sperm analysis (CASA) that has been shown to be an objective tool for measuring sperm characteristics (Elofsson, Van look, Borg, & Mayer, 2003; Kime et al., 1996; 2001; Rurangwa, Kime, Ollevier, & Nash, 2004). The image analyser was set as follows: frame rate 50 Hz; no. of frames 25; minimum contrast 11. To avoid including measurements taken of sperm cells moving due to drift or Brownian movement, threshold values for the only two optional settings defining static cells, that is, VAP and VSL, were set at 10 µm/s. The same method has successfully been used in previous studies (e.g., Janhunen et al., 2009; Liljedal et al., 2008; Urbach et al., 2005). The parameters assessed were average path velocity (VAP), straight-line velocity (VSL), and curvilinear velocity (VCL) (Rurangwa et al., 2004). The data from the CASA revealed that these three sperm velocity parameters measured at 10 s after activation were significantly correlated, and Pearson’s correlation coefficient was 0.96, 0.87, and 0.79 when comparing VCL and VAP, VSL and VAP, and VCL and VSL, respectively (N = 32 and p < 0.001 in each of the three correlations). Because of a lack of target (an egg) and no gradient in the concentrations of OF, we could not assume a straight-line swimming behavior of sperm cells. Based on this, and the high correlation between the different velocity measurements (see above), we chose to use curvilinear velocity (from now on termed “sperm velocity” or “VCL”), as the measure of sperm velocity in our statistical models. Velocity estimates for each female–male (OF-ejaculate) combination correspond to the mean velocity of all motile cells analyzed. We did not add any artificial substances, for example sperm extenders, to the milt.

2.4 Handling of female gametes

The stripped eggs embedded in OF from each female were stored in the dark at approximate lake water temperature until handling. In each block of the fertilization experiment (see Table 1), eggs from two females and OF from two females were treated in three different ways before sperm was added and eggs fertilized. We started by distributing eggs from each of the two females in each block into six batches (into six petri dishes) with three different treatments each with two replicates of similar egg numbers (similar egg numbers in each petri dish from one particular female, but the number varied between the females with minimum and maximum number of eggs per petri dish being 27 and 170, respectively) and OF volume. Within two of the six batches, the eggs were physically separated from the OF. The petri dish containing the eggs and OF was first tilted at approximately 30° long enough for the OF to drain from most of the eggs (1–2 min). The OF, now located at the lower part of the petri dish, was carefully removed with a pipette from the lowest point of the petri dish. This way of separating eggs and OF, by draining OF from the eggs, is probably similar to the draining of OF as reported by Lehnert et al. (2018). Thereafter, the OF was returned to the same eggs (i.e., “own OF”). In the next two batches, the eggs and OF were separated as described above, but the OF was not returned to the same eggs; rather, the eggs were mixed with OF from the other female in the block (i.e., “foreign OF”). In the remaining two batches,
the eggs and OF were not separated (i.e., "control"). The average volume of OF removed was 2.9 ml (SD = 1.58 ml), which gives approximately 0.5 ml OF per replicate. We did not wash the egg before sowing OF, nor did we pat the eggs dry or add any other artificial substances to the eggs or OF as we do not know the unforeseen effects of such treatment. The time elapsed from OF was removed from the eggs until it was re-added or exchanged was approximately 45 min.

2.5 | Fertilizations and rearing

Using micropipettes, milt from the two males was first added to the bottom of a glass jar carefully controlling for not allowing physical contact between the ejaculates. To ensure that differences in fertilization success between each of the two competing males in a pair were independent of initial difference in sperm numbers, the volume of the milt used in the sperm competition trials was adjusted according to the spermatocrit values to give an approximately equal number of sperm cells from each male. Fertilizations was conducted by adding 50 ml of lake water, mixing the two milt samples, and then gently pouring this mix over to a 500-ml plastic jar already containing eggs embedded in OF. This was followed by 5 s gentle movement of the jar containing all gonadal products. The ratio of OF to water was approximately 1 to 100 during these first 5 s. Thereafter, the 500-ml plastic jar was filled with water, in order to dilute sperm concentrations and avoid polyspermy, and sealed. Each of the batches was then stored separately in a refrigerator at 4°C–6°C until transport to the hatchery, which occurred within 24 hr after fertilization. The eggs from each of the fertilizations were randomly positioned in two separate tanks and kept under a natural light regime in the hatchery at the University of Tromsø for the next 60 days (from fertilizations on 25 September in both 2011 and 2012) until they were killed with 75% ethanol when the experiment was terminated (the 15 November in both years). Untreated 6°C water was constantly exchanged in the tanks during this period.

2.6 | Genotyping

Offspring and parents were genotyped using microsatellite DNA analysis. DNA extraction was done with the MasterPure™ DNA and RNA Purification Kit (EPICENTRE Biotechnologies, Cat. No. MC85200), following the instructions for tissue samples under point B user manual "Purification of Total Nucleic Acids and Precipitation of Total DNA." When optimizing quantities of DNA, we used the Quant-iT™ dsDNA Broad-Range Assay Kit (Invitrogen Detection Technologies, Lot. 1116429). Paternity was assigned using microsatellites and PCR. The PCR products (5 μl total volume) contained 200 ng DNA, 20 mM of primer forward, 20 mM primer reverse, and 1.25 units of AmpliTaq Gold 360 MM (Applied Biosystems, Life Technologies). The contents of each well were mixed by careful pipetting, no vortex. The plate was then sealed, spun shortly, and subjected to thermal cycling. PCR was performed with a Veriti™ 96-Well Thermal Cycler (Applied Biosystems, Life Technologies) by the following steps: Stage 1 (five cycles), 95°C 1:00 min, 55°C 0:20 min, 72°C 0:25 min; Stage 2 (five cycles), 95°C 0:30 min, 55°C 0:20 min, 72°C 0:25 min; Stage 3 (28 cycles), 95°C for 0:20 min, 55°C for 0:20 min, 72°C for 0:25 min; and Stage 4 (one time only), 72°C 20:00 min, 15°C for final temperature. The PCR products were then kept at 4°C until further handling. PCR products were separated on a 3500xl capillary sequencer (Applied Biosystems, Foster City, CA, USA), and alleles were identified by the ABI GeneMapper Software, version 4.1 (Applied Biosystems, Life Technologies). We used the following three microsatellite loci for genotyping: Smm_22 and Smm_24 (Crane et al., 2004), and Omm_1070 (Rexroad et al., 2001). In the present study, the number of unique alleles of Smm_22, Smm_24, and Omm_1070 of the 24 males and 24 females (parents) producing the 72 different fertilizations was 11, 10, and 4, respectively. The same microsatellites have also previously been reported as highly polymorphic in the present char population (see Table 1 in Westgaard, Klemetsen, & Knudsen, 2004). The offspring were always identified as the offspring of one of the two fathers involved in each sperm competition, and one microsatellite was in most cases enough to unambiguously assign offspring to one father.

2.7 | Statistical analyses

Statistical analyses were carried out similar to Evans et al. (2013) by using R version 3.3.2 (R Core Team, 2016). Two statistical analyses were conducted: sperm velocity analysis and paternity analysis. In the sperm velocity analysis, we tested potential effects of male identity and OF identity on sperm velocity (VCL). In this analysis, VCL was measured at 10 s after activation for the 32 different levels of the interaction variable between male identity and OF identity, and each measurement was replicated twice. Thus, the data set for the sperm velocity analysis contains 64 observations. We used linear mixed models (lmer) from the lme4 package (Bates, Maechler, Bolker, & Walker, 2016) including sperm velocity VCL (see above) as a continuous response variable and OF identity, male identity, and their interaction as random effects. Models with and without a specific random factor were compared using likelihood ratio tests from the R package lme4test (Zeileis & Hothorn, 2002), starting from the full model including the interactions.

In the paternity analysis, we tested the effects of both male identity and egg-donor (i.e., the identity of the female’s eggs as opposed to identity of the female’s OF) on the paternity success of the focal male in the different fertilization treatments. The data set of this analysis consisted of 72 observations (i.e., in vitro fertilizations) with on average nine offspring, as described above. We used a binomial generalized linear mixed models (glmer from lme4) with a two-column matrix as the response variable, where the first column consists of the numbers of offspring sired by the focal male, and the second column consists of the numbers of offspring sired by the second male. Note that this is equivalent to specifying relative paternity as the response variable together with specifying the parameter “weights” as the vector of the number of offspring sired by both males. Predictors were the treatments...
TABLE 2 Results from linear mixed models testing random effects of male identity (male ID), identity of ovarian fluid (OF ID), and their interaction, on the response variable sperm velocity in our data set of N = 64 observations. Sperm velocity was measured in ovarian fluid (OF) diluted in water. The standard deviations of the random effects were estimated by using the function lmer in the R package lme4, while the chi-square statistics and the p-values were measured by comparing models with and without this factor by likelihood ratio tests using the R function anova.

| Source              | SD    | Chi-square | p-Value |
|---------------------|-------|------------|---------|
| Male ID             | 14.69 | 17.104     | <0.0001 |
| OF ID               | 17.89 | 25.73      | <0.0001 |
| OF ID: male ID      | 5.56  | 0.395      | 0.53    |

The study was carried out in accordance with ethical guidelines stated by the Norwegian Ministry of Agriculture and Food through the Animal Welfare Act. According to these guidelines, we were not supposed to—and therefore do not—have a specific approval or approval number.

3 | RESULTS

3.1 | Sperm velocity analysis

Sperm velocity in OF was significantly influenced by both male identity and OF identity (Table 2). That is, sperm from some males swam generally faster than sperm from other males in OF, and OF from some females affected sperm speed more than OF from other females. On the other hand, no interaction effect was revealed between OF identity and male identity (Table 2), suggesting that OF did not affect sperm velocity according to individual characteristics of ejaculates.

3.2 | Paternity analysis

There was no support for OF affecting relative paternity between the two competing males. That is, relative paternity, measured as the ratio between the number of offspring sired by the focal male and the number of offspring sired by both males, was not influenced by our experimental exchange of OF between “egg batches” (see Table 3). Figure 2 illustrates that the relative paternity was not influenced by our experimental exchange of OF. Yet, relative paternity tended to differ between the treatments “control” versus “own OF” (p = 0.055, Table 3). However, this tendency was absent when comparing “own” versus “foreign OF” (p = 0.334) and when comparing “foreign” versus “control” (p = 0.348). Male identity significantly affected relative paternity (Table 3), but egg-donor had no effect on paternity. Thus, although OF identity affected the sperm velocity (first analysis), there was no effect of any female parameter (egg-donor or OF) on the relative paternity.

Relative velocity of the sperm from the two competing males did not affect relative paternity (Figure 1), suggesting that the observed difference in swimming speed between the two males in a pair was not important for fertilization success under our experimental setup.

TABLE 3 Results from the generalized linear mixed models testing the effects of ovarian fluid (OF) on relative paternity. The response variable, relative paternity, was measured as number of offspring sired by the focal male divided by the sum of offspring sired by both males in each of the 72 in vitro fertilizations. Relative sperm velocity (i.e., the sperm velocity of the focal male minus sperm velocity of the competing male) was included as a continuous covariate. The treatments (“control,” “own OF,” and “foreign OF”) are dummy variables. In the table below, the last mentioned treatment on a line is the base value in this comparison. For example, “Treatment ‘control’ versus ‘own OF’” means that the effect of treatment “control” is estimated when treatment “own OF” is the base value. The test of “own versus control” (first analysis), there was no effect of any female parameter (egg-donor or OF) on the relative paternity.

| Source            | Estimate | SE   | z-Value | p-Value |
|-------------------|----------|------|---------|---------|
| Intercept         | 0.305    | 0.807| 0.378   | 0.706   |
| Treatment “control” versus ‘own OF’ | −0.507 | 0.264| 1.921   | 0.055   |
| Treatment “own OF” versus “foreign OF” | 0.266 | 0.275| 0.966   | 0.334   |
| Treatment “foreign OF” versus “control” | 0.241 | 0.257| 0.939   | 0.348   |
| Relative sperm velocity | −0.006 | 0.004| −1.281  | 0.2     |

| Source          | Estimate | Var | Chi-square | p-Value |
|-----------------|----------|-----|------------|---------|
| Egg-donor ID    | −0       | −0  | >0.999     |         |
| Male ID         | 4.887    | 13.204| 0.0003    |         |
| Egg-donor ID: male ID | −0 | −0 | >0.999 |         |
3.3 | Power analysis

As we did not find any significant fixed effects in the paternity analysis, we performed a power analysis by using the R package simr (Green & MacLeod, 2016). When using an effect size approximately at the level of the estimated coefficients (0.25 and 0.50, see “Estimate” in Table 3), the power of our tests of the influence of OF on relative paternity is indeed very low (0.07 and 0.18). However, statistical power is a function of the effect size, which in our experiment is the difference in relative sperm velocity between the two males in all observations with this value of relative sperm velocity, divided by the total number of offspring sired by both males in all observations with this value of relative sperm velocity. Note that the points represent different numbers of observations: six observations in the full blocks, and two or four observations in the half blocks.

4 | DISCUSSION

Our main finding is that exchange of OF between egg batches from different females did not affect the males’ relative fertilization...
success under sperm competition. Although sperm from some males generally swam faster in OF than sperm from other males, differences in sperm velocity between males had no effect on their relative fertilization success. Ovarian fluid from some females increased sperm speed more than OF from other females, but OF did not affect sperm velocity according to individual characteristics of ejaculates. Additionally, there was no indication that the eggs themselves favored sperm from one male over the other under sperm competition. Thus, the only two studies which have intraspecifically exchanged OF between eggs so far conclude that the effect size of cryptic choice exerted by OF is either small or absent (Evans et al., 2013; the present study). This result concurs to the main conclusion in a recent review, which found few clear demonstrations of cryptic female choice (Firman et al., 2017).

Common for experimental studies, our results may be affected by the applied methodology. First, the lack of effect when exchanging different OFs on paternity could result from our inability to remove all OF from the eggs. That is, we were not able to remove the last remains of the OF bound to the egg surface and the micropyle. On the other hand, we cannot exclude the possibility that physically removing the last remains of the original OF from the eggs by washing the eggs with water-based isotonic fluid (Evans et al., 2013; Yeates et al., 2013) could affect the eggs and consequently the result of sperm competition. Evans et al. (2013) reported no effect of exchanging OF between females after washing the eggs with an artificial ovarian solution before adding foreign OF. Thus, the consistent results between our two studies—using slightly different methods on taxonomically closely related species—suggest that OF is not a medium exerting strong cryptic selection of sperm. It may be advocated that both methods applied to remove OF from the eggs are flawed, but this does not concur with the expected effect of OF on paternity documented interspecifically when eggs were also washed by isotonic solution (Yeates et al., 2013). It is therefore unlikely that different methods of separating OF from ova in the two studies caused the different results. Second, sperm velocity measurements were initiated 10 s after activation of sperm cells in the present study. Although as much as 80% of fertilizations in the river spawning sockeye salmon (O. nerka) may occur within 5 s (Hoyersak & Liley, 2001), charr spawn in still water and show male–female interaction effects when sperm swim in OF as late as 30 s after activation. Our sampling delay is shorter or comparable to previous studies using model species that do not spawn in still water (Alonzo, Stiver, & Marsh-Rollo, 2016; Evans et al., 2013; Yeates et al., 2013). Third, the concentration of OF (diluted in water) was 1% during the fertilizations in this study. Yeates et al. (2013) used the same (1%) concentration and reported significant effects of OF on paternity under interspecific sperm competition trials, whereas no effect of OF was revealed from intraspecific fertilizations at 10% solutions (Evans et al., 2013). Spawning behavior of charr in the present population has been studied by Sørum et al. (2011) and Brattli et al. (2018). The studies show that (a) more than 50% of the spawning events occur under sperm competition, (b) mean number of males is 2.9 s at egg release and increases to more than four males within the next 2.0 s, (c) the first male to release milt ejaculated from 0.15 s before to 1.9 s after the eggs are shed, and (d) the average time delay in gamete release under sperm competition between the first and the subsequent males is estimated as 0.68 s (Sørum et al., 2011). Moreover, some males have the advantage of spawning physically relatively close to the spawning female, whereas the remaining males spawn further away (Brattli, Egeland, Nordeide, & Folstad, 2018; Sørum et al., 2011). The exact concentration of OF diluted in water at the time of fertilizations in natural spawning events is not known for this population, but most likely it varies a lot between fertilizations and competing males. Thus, we cannot conclude whether 1% or 10% OF to water mimics the natural spawning conditions more closely. Fourth, a growth period from fertilization to sampling is needed for DNA-sampling in order to estimate paternity. In the present study, this period lasted 60 days, that is, the same as that of Yeates et al. (2013), compared to 28 days in Evans et al. (2013). It seems inevitable that some eggs do not develop during this period as some eggs may be unfertilized and some zygotes malformed or dead. It is however unknown what causes specific mortality at this early stage. Yet, one possibility is that specific mortality differs because of varying “egg quality” due to physiological nonoptimal timing or other conditions during the artificial stripping and handling of gametes (Bobé & Labbé, 2010; Lahnsteiner, Weismann, & Patzner, 1999). An alternative explanation is that one of the two males in a block sired offspring with higher survival than the other male, that is, due to genetically superiority (Evans et al., 2013; García-González, 2008). Under such “good-sperm effects,” one would expect a positive and significant association within pairs of males between mortality of the eggs on the one hand and skewness in paternity on the other hand. A post hoc test carried out on the present data showed no association between relative paternity of the males and the proportion of eggs surviving ($r_s = 0.134, p = 0.44$, $N = 36$, Spearman’s correlation coefficient after pooling both replicates). This result concurs with those previously reported from two independent studies using individuals from the same charr population, similar experimental designs, and the same rearing equipment and methods as in the present study (Egeland et al., 2015; Liljedal et al., 2008). It is therefore unlikely that the actual fertilization success we measure is caused by differential mortality or different developmental ability of the embryos sired by the two males in each block (see García-González, 2008 for further discussion). Fifth, sperm velocity had no significant effect on relative paternity in our study (Table 3). The two males in each block were picked at random from the spawning grounds, and this lead to small within-pair differences in sperm velocity. That is, relative sperm velocity differed by less than 10% in 26 of the 48 measurements (with “foreign OF,” “own OF,” and “control” combined). Such small between-males differences in sperm velocity of our experimental pairs might explain why we did not find significant effects of relative sperm velocity on paternity. This is contrary to Egeland et al. (2015) who reported relative sperm velocity and motility as the best predictor of male fertilization success in charr from our study population. In the latter study, the two males whose sperm were competing to
fertilize eggs in vitro were caged together for four days prior to ferti-
zizations in order to deliberately produce one dominant male with
low sperm velocity and one subordinate male with high sperm veloc-
ity (see also Egeland et al., 2016; Rudolfsen et al., 2006). This “pro-
duction” of large differences in sperm velocity might explain the
contrasting effects of sperm velocity on paternity in the two studies.
In accordance with this explanation, Evans et al., (2013) who at-
tributed chinook salmon males’ relative paternity to variation in the
relative sperm competitive ability reported large variation in relative
sperm velocity between the focal and the competing chinook males
in their sperm competition experiment (fig. 3 in Evans et al., 2013).
Sixth, a total of 649 offspring were genotyped in this study (see
Materials and methods and Table 1), which is about 1/3 of the 1937
offspring genotyped by, for example, Evans et al., (2013). Moreover,
our experiment consisted of four full blocks (one block consists of
two trials) in addition to four trials (“half blocks”), which is approxi-
mately half the number of blocks (or trials) compared to Evans et al.,
(2013). The power analysis suggests that our sample size is large
enough to detect medium-sized effect sizes (changes in relative pa-
ternity from each male in the sperm competitions carried out in OF
from two females), but not large enough to detect the small changes
in relative paternity found in our study. Seventh, we picked the charr
in each block by random and hence cannot exclude the possibility
that the two females in each block (or trial) have very similar OF, for
example, due to being closely related. If so, we should expect no ef-
fect on relative paternity. Eight, as the only study of this kind so far
we included a second control group (“own OF,” see Materials and
methods), to test for the potential effect of our handling of the fe-
male reproductive products by removing and then re‐adding the
of sperm:egg ratios during fertilizations in Arctic charr.

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CONFLICT OF INTEREST
None declared.

AUTHORS’ CONTRIBUTION
SAK, JTN, GR, LF, and IF designed and performed the experiment.
SAK and GR did the sperm analyses. SAK did the paternity analysis
and IF reared the eggs. BL, KR, and JTN did the statistical analyses.
All authors took part in the interpretation of the results and wrote
parts of the manuscript, although most of the writings was carried
out by SAK, JTN, and IF. All authors have read and approved the final
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DATA ACCESSIBILITY
The data used in the Sperm velocity analysis and Paternity analysis
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