See-Thru-Gonad zebrafish line: developmental and functional validation

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Abstract

Zebrafish are an important model species in developmental biology. However, their potential in reproductive biology research has yet to be realized. In this study, we established See-Thru-Gonad zebrafish, a transparent line with fluorescently labeled germ cells visible throughout the life cycle, validated its gonadal development features, and demonstrated its applicability by performing a targeted gene knockdown experiment using vivo-morpholinos (VMOs). To establish the line, we crossed the zf45Tg and mitfa^{w2/w2}, mpv{17}^{6A/B} zebrafish lines. We documented the in vivo visibility of the germline-specific fluorescent signal throughout development, from gametes through embryonic and juvenile stages up to sexual maturity, and validated gonadal development with histology.

We performed targeted gene knockdown of the microRNA (miRNA) miR-92a-3p through injection of VMOs directly to maturing ovaries. After the treatment, zebrafish were bred naturally. Embryos from miR-92a-3p knockdown ovaries had a significant reduction in relative miR-92a-3p expression and a higher percentage of developmental arrest at the 1-cell stage as compared with 5-base mismatch-treated controls. The experiment demonstrates that See-Thru-Gonad line can be successfully used for vertical transmission of the effects of targeted gene knockdown in ovaries into their offspring.

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Introduction

Zebrafish (Danio rerio) have undergone a remarkable journey from their origin in the rivers of India and Pakistan to the forefront of science. There are many advantages of zebrafish research including relatively low costs, short generational time, rapid embryonic development, large transparent embryos and a general robustness allowing for manipulations (Veldman & Lin 2008). Despite this, one area of research for which the potential of zebrafish has yet to be realized is reproductive biology.

Several lines have been established in an attempt to facilitate research on reproduction in zebrafish. The zf45Tg line was created in Lisbeth Olsen's lab using the vasa regulatory regions fused with enhanced green fluorescent protein (Tg(vasa:vasa-eGFP) construct) to fluorescently label cells of the germline (Krøvel & Olsen 2002). vasa, a conserved gene coding for a DEAD box RNA helicase, is the best known germ cell marker, with conserved and specific expression in the germline across metazoans (Gustafson & Wessel 2010). The zf45Tg line successfully allows the tracking of primordial germ cells (PGCs) from approximately 24 h post fertilization (hpf) onward (Krøvel & Olsen 2002). However, additional studies have shown that only maternal GFP signal is detected in the germline during early development, and that zygotic transcription does not begin until after sexual differentiation when the germ cells have entered meiosis around 21 days post fertilization (dpf) (Krøvel & Olsen 2004, Leu & Draper 2010). The Tg(bactin:eGFP) line was created to observe sex differentiation in zebrafish (Hsiao & Tsai 2003). beta actin (bactin) is ubiquitously expressed throughout zebrafish but its expression is significantly higher in ovaries, allowing their identification by the contrast in signal intensity (Hsiao & Tsai 2003). However, the Tg(bactin:eGFP) line cannot be used to identify gonads before 14 dpf, or testis at any stage (Hsiao & Tsai 2003). More recently, the construct comprising promoter elements for ziwi (piwil1), the zebrafish homolog for Drosophila piwi, and eGFP, was used to create the uc1Tg line (Leu & Draper 2010). ziwi, like vasa, is specifically expressed in the germline of zebrafish throughout development (Draper et al. 2007, Houwing et al. 2007). In this transgenic line, ziwi:eGFP is maternally supplied in embryos, while zygotic GFP can be first detected around 7 dpf (Leu & Draper 2010). Therefore, the line is less dependent on maternal eGFP to label germ cells early in development than the zf45Tg line. However, after hatching, the usefulness of these lines in larval and subsequent development stages is limited in vivo due to interference from pigment and the general opaqueness
of zebrafish tissue. While eGFP signal can be detected, it only allows for a rough comparison of fluorescence intensity, and does not allow for observation of gonadal structure without excision of the gonad.

In wild-type zebrafish, the pigmentation pattern consists of dark blue stripes composed of three types of pigment cells: black melanophores, yellow xanthophores and reflective iridophores (Rawls et al. 2001, White et al. 2008, Krauss et al. 2013). The tissue opaqueness can be reduced by chemical treatment, such as treating zebrafish embryos with 1-phenyl 2-thiourea before the initiation of melanogenesis (Karlsson et al. 2001). However, there is an issue with fish welfare since fish treated with 1-phenyl 2-thiourea have a marked increase in anxiety during experimental trials (Parker et al. 2013). Alternatively, some mutants such as the mtf1aw2/w2; mpv17b18/b18 show transparent phenotypes. This line, which phenotypically resembles casper (roy −/−; nacre −/−), is a double mutant for the nacre and transparent (tra) phenotypes (White et al. 2008). The nacre phenotype is caused by a polymorphism in microphthalmia-associated transcription factor alpha (mitfa), which results in a premature stop codon (Lister et al. 1999). Zebrafish with mutant mitfa lack melanocytes completely and compensate it with an increased number of iridophores (Lister et al. 1999, White et al. 2008). The transparent phenotype is caused by a mutation in mitochondrial inner membrane protein 17 (mpv17) (Krauss et al. 2013). The mpv17 transcript contains a splicing error between exons 2 and 3, which causes the deletion of 19 nucleotides from the open reading frame and ultimately results in a frameshift and early stop codon (Krauss et al. 2013). Transparent (−/−) zebrafish have a reduced number of iridophores and melanocytes, uniformly black pigmented eyes and a translucency of the skin (Krauss et al. 2013).

The Tg(bactin:eGFP) line was recently crossed with roy (−/−) and albino (alb; −/−) mutants to create β-ruby, a transparent and eGFP-expressing zebrafish line for use in reproductive biology (Akhter et al. 2016). While the line provides improved transparency and visualization of the ovary it contains the same limitations as the parent Tg(bactin:eGFP) line; gonads cannot be identified until after 14 dpf, and differentiated testis do not express eGFP signal.

miRNAs are short nonprotein-coding RNA molecules with a primary function in posttranscriptional repression of messenger RNAs (mRNAs) through associating with the miRNA-inducing silencing complex (miRISC) and then binding to the 3′ untranslated region of an mRNA (Bartel 2009, Fabian & Sonenberg 2012). Depending on the complementarity between the miRNA and mRNA, the mRNA may be sequestered or degraded (Fabian & Sonenberg 2012, Bizuayehu & Babiat 2014). The role of miRNA during the gonadal development in fish is still largely unknown (Bizuayehu & Babiat 2014). Several recent studies have reported on miRNA expression in teleost gonads (Bizuayehu et al. 2012, Jing et al. 2014, Xiao et al. 2014, Vaz et al. 2015, Juanchich et al. 2016, Tao et al. 2016). However, miRNA expression during gonadal development of zebrafish, a primary teleost model, has not been profiled yet due to the small size and the challenges associated with dissecting gonads in juvenile stages.

In vivo methods for functional analysis of miRNA in zebrafish gonad have yet to be developed. Recently, targeted gene knockdown by injection of morpholino oligonucleotides antisense for an miRNA precursor was developed for the zebrafish brain (Kizil et al. 2013). There is potential that a zebrafish line optimized for reproductive biology could allow this technique to be adapted for gonadal studies as well.

The first objective of our study was to establish a stable line for use in reproductive biology with visible and fluorescently labeled germ cells throughout the zebrafish life cycle. We crossed the transgenic zf45Tg and mtf1aw2/w2; mpv17b18/b18 double mutant lines to create the See-Thru-Gonad line. We validated its functionality throughout gonadal development by a combination of in vivo imaging and histology. The second objective was to demonstrate the applicability of See-Thru-Gonad for miRNA functional studies. We performed targeted gene knockdown of the miRNA miR-92a-3p in sexually maturing ovaries and quantified its effects in embryos originating from the treated ovaries. miR-92a-3p was chosen for the trial as it had been shown previously to be abundant in both zebrafish gonads and embryos (Vaz et al. 2015) and had been linked to ovarian development in mammals (Liu et al. 2014, Maalouf et al. 2015); but its function had yet to be explored in teleost gonads. We demonstrated that miR-92a-3p was maternally inherited and that the effect of targeted gene knockdown is vertically transmitted to the next generation.

Methods

Fish

All husbandry and experimental procedures were performed in accordance with the Norwegian Regulation on Animal Experimentation (The Norwegian Animal Protection Act, No. 73 of 20 December 1974) and were approved by the National Animal Research Authority (Utväg for forsk med dyr, frosøksdyrutvalget, Norway) General License for Fish Maintenance and Breeding (Godkjenning av aveling for frosøksdyr) no. 17.

Zebrafish were maintained in an Aquatic Habitats recirculating system (Pentair, Apopka, FL, USA) following standard zebrafish procedures (Westerfield 2000). Zebrafish diet consisted of a daily mix of newly hatched Artemia nauplii (Pentair) and SDS zebrafish-specific diet (Special Diet Services, Essex, UK) following the manufacturers’ recommended feeding regime.
**Creation of the See-Thru-Gonad line**

The parent lines used were the zf45Tg line (Krovel & Olsen 2002) and the mitta\(^{w2}/w2\), mpv17\(^{b18}/b18\) double mutant line, both obtained from The Norwegian Zebrafish Platform (zebrafish.no), Norwegian University of Life Sciences (Oslo, Norway). Initially, ten F\(_1\) breeding pairs of each line were spawned naturally with males of the mitta\(^{w2}/w2\), mpv17\(^{b18}/b18\) mutant line, as only maternal eGFP signal is visible in Tg(vasa:vasa-eGFP) embryos (Krovel & Olsen 2002, 2004). At 2 dpf, the F\(_1\) embryos were screened and 100% expressed the GFP signal. However, as nacre and transparent are recessive mutations, all F\(_1\) individuals were pigmented. Upon reaching sexual maturation, the F\(_1\) fish were intercrossed, producing 3000 F\(_2\) embryos. The embryos were screened at 2 dpf for lack of melanophores and presence of eGFP signal (Fig. 1A). The transparent mutation is not visible in embryos; therefore, the remaining fish were raised until 6 weeks post fertilization (wpf), at which time the homozygous transparent mutants could be visually identified and collected (Fig. 1B). The fish showing all three desired phenotypes were intercrossed to create the F\(_3\) generation.

The F\(_3\) were screened as before for the GFP signal (Fig. 1A). Because the F\(_2\) fish were heterozygotic for Tg(vasa:vasa-eGFP), an estimated 75% of them were expected to express the signal. GFP-expressing fish were further reared until sexual maturation, at which time they could be screened for homozygosity of the Tg(vasa:vasa-eGFP). Females were crossed with AB line males, and their offspring screened for GFP as before. Fish whose offspring were all expressing GFP were considered homozygous, whereas fish whose offspring were mixed were considered heterozygous (Fig. 1C). In the case of males, the transgene phenotype was observed directly in sperm (Fig. 1C). Homozygotic F\(_3\) fish were considered as a stable See-Thru-Gonad line, and their offspring were used for subsequent experiments.

**DNA and RNA isolation**

For DNA isolation, three-month-old See-Thru-Gonad fish were anesthetized using 50 mg/L MS-222 (Tricaine; Sigma Aldrich) buffered with equal parts of sodium bicarbonate (NaHCO\(_3\)) before caudal fin clips were taken. DNA was immediately extracted using the QuickExtract DNA Extraction Solution (Illumina, San Diego, CA, USA) according to the manufacturer's protocol.

For RNA extraction, four fish (two males and two females) were first killed with 200 mg/L buffered MS-222 followed by decapitation. Fast and slow muscle as well as skin from the trunk were excised and RNA was immediately extracted using QiAzoL Lysis Reagent (Qiagen) following the manufacturer's instructions. Embryos were collected directly to QIAzol. RNA quality was assessed using 1% (w/v) gel electrophoresis and quantified using a NanoDrop ND-1000 (Thermo Fisher Scientific, Saven & Werner AS, Kristiansand, Norway). Total RNA was reverse transcribed using the QuantiText Reverse Transcription Kit (Qiagen) following the manufacturer's protocol.

**Sequencing**

To obtain genomic fragments of mpv17, mitfa, and the vasa:vasa-eGFP transgene, gene-specific primers were designed or taken from literature (Supplementary Table 1, see section on supplementary data given at the end of this article). For mpv17, primers were designed to amplify a CDNA fragment as well. Polymerase chain reaction (PCR) was performed to amplify the target genes using LongAmp Taq DNA polymerase (New England BioLabs, Ipswich, MA, USA). The PCR products were excised from a 1% (w/v) agarose gel and eluted using the QIAquick gel extraction kit (Qiagen). Amplified fragments were ligated into the pCR4-TOPO vector (Thermo Fisher Scientific) and transformed into OneShot chemically competent E. coli cells. The plasmid DNA was extracted and purified using the QiAprep Miniprep kit (Qiagen) and sequenced in both directions.
directions using M13 primers with the Big Dye Terminator 3.1 (Applied Biosystems) sequencing template preparation method. Sequencing reactions were analyzed by the DNA Sequencing Lab at the University of Tromsø, Norway.

**Visualization of germline in vivo**

Cells of the germline were identified by their expression of the *vasa:vasa-eGFP* transgene (Krøvel & Olsen 2002). Embryos at 2 dpf were imaged directly, while larvae at 4 dpf, as well as fish at 2, 3, 6, 9 and 12 wpf were anesthetized as described previously before in vivo imaging. Because many zebrafish diets may cause nonspecific fluorescent signal within the digestive tract, all fish at 2 wpf and older were fasted for a minimum of 24 h before imaging. For gonad dissection, the zebrafish were killed as described previously and the gonad was removed under epifluorescent light using fine dissecting tools. For spermatozoa collection, mature males were anesthetized and patted dry with tissue paper before gently stroking the side of the fish. Spermatozoa were collected from the genital pore using a 10-µL pipette with gentle suction and diluted in 50 µL Hanks’ balanced salt solution (Sigma Aldrich). All visualization of the gonads was performed using an AxioZoom V.16 microscope (Carl Zeiss) equipped with an AxioCam MRm monochrome camera (Carl Zeiss) and Zen Pro (2012; Carl Zeiss) imaging software. Images were taken using either bright field or epifluorescent light with an enhanced green fluorescent protein filter (excitation: 488 nm, emission: 509 nm).

**Gonadal histology**

The gonads of 3, 6, 9 and 12 wpf zebrafish were examined. After being killed, the fish were truncated and the body cavity was opened posterior to the pectoral fins before being fixed with Bouin’s solution (Sigma Aldrich) overnight at 4°C. After fixation, the samples were embedded in paraffin, sectioned and stained using hematoxylin and eosin as described previously (Presslauer et al. 2014). At 3 wpf, before sexual differentiation, undifferentiated gonads from three individuals (n = 3) were examined. From 6 wpf onward, four fish of each sex (n = 4 + 4) were examined at each time point. Staging of gonads was performed according to the previously established literature (Selman et al. 1993, Maack & Segner 2003, Neumann et al. 2011).

**Knockdown of miRNA in adult zebrafish**

A VMO complementary to the dre-miR-92a-3p guide strand (5’-TACGGCCGGACAGTGCATTACCC-3’) was well as a 5-base mismatch VMO (5’-TTACGGCCGGACAGTGCATTACCC-3’; underlined letters indicate the mismatches) as a negative control were designed and produced by Gene Tools LLC (Philomath, OR, USA). The treatment was administered to 12 wpf females (Supplementary Table 2). For each injection, fish were anesthetized and injected directly to the posterior portion of the ovary using a 10.0 µL syringe equipped with a 34 g needle (Hamilton, Bonaduz, Switzerland). Injections were either 2.0 or 3.0 µL of 0.5 mM morpholino or the mismatch control.

After 3 days of recovery, the fish were allowed to spawn naturally. From each spawning, fifteen embryos were collected and RNA was immediately extracted as described previously. The remaining embryos were monitored and developmental features were recorded at 4 and 24 hpf.

**Real-time quantitative RT-PCR**

Two reference genes (miR-26a-5p and miR-99a-5p) were chosen based on their lack of similarity to the targeted miR-92a-3p and their relatively high abundance in the maturing zebrafish ovary. TaqMan MicroRNA Assays (Thermo Fisher Scientific) were performed for both reference and target genes on a LightCycler 480 (Roche, Mannheim, Germany). The thermal cycle conditions used were 95°C for 10 min, followed by 55 cycles of 95°C for 15 s, 60°C for 1 min and 72°C for 1 s. The 20 µL PCR consisted of 1× TaqMan small RNA assay, 30× diluted reverse transcription product and 1× TaqMan Universal PCR Master Mix II. All samples were run in duplicate with no reverse transcription and no template negative controls. Cycle threshold (Ct) values were determined using the LightCycler 480 software cycle cut-off value of 40. A five-point serial dilution curve (dilutions 1:15–1:240) was used to determine the efficiency of the PCR processes. Normalization between samples was performed using miR-26a-5p and miR-99a-3p as reference genes and normalization factors were calculated using geNorm (Vandesompele et al. 2002), which provided a stability value of 0.863.

**Statistical analyses**

Percentage data (frequency of embryos with arrested development) were arcsine square roots transformed. The effect of treatment at two injection doses on miR-92a-3p expression and occurrence of developmental arrest in early embryos was evaluated using a two-way ANOVA. Since no significant first-order interaction treatment × dose was found, the model was broken down and each injection dose was analyzed separately with a simple ANOVA. The effect was considered significant at P value of 0.05 or less (Zar 1999).

**Results**

**Identification of See-Thru-Gonad mutations**

Sequencing of zebrafish *mpv17* cDNA confirmed the deletion of 19 nucleotides within the coding sequence, resulting in a frameshift and early stop codon (Supplementary Fig. 1A) and was consistent with a previous report (Krauss et al. 2013). PCR for the corresponding genomic region using conditions optimized for an increased product size amplified a region approximately 3 kb in length, compared with an expected amplicon of 284 nucleotides (Supplementary Fig. 1B). Sequencing of this fragment from both ends revealed an insert of approximately 3000 bp near the end of the second exon in the *mpv17* gene (Supplementary Fig. 1C, Supplementary Table 3). Sequencing of zebrafish *mitfa* confirmed the polymorphism reported...
previously within exon 7 (Lister et al. 1999). This polymorphism resulted in a UAA stop codon, resulting in premature termination of translation (Supplementary Fig. 1D). In addition, the vasa:vasa-eGFP transgene was successfully amplified (Supplementary Table 3).

**Visualization of the germline in vivo**

At 2 dpf, the See-Thru-Gonad line showed enhanced resolution of germ cells when compared with the zf45Tg line (Fig. 2). A comparison of the lateral and dorsal views shows most PGCs to be on the same lateral plane in small clusters. When counting PGCs, an average of 26 (s.d. = 3.7, n = 5) could be identified in the See-Thru-Gonad line, whereas a lack of transparency made identification of individual PGCs in the zf45Tg line not possible.

At 4 dpf, germ cells were clearly visible near the gonadal anlagen (Fig. 3A). A lateral view showed the germ cells on the same lateral plane, while a dorsal view allowed observation of the gonadal structure at this time. At 2 wpf, the gonad was not identifiable using transmitted light, but imaging using epifluorescent light identified a compact gonad approximately 0.5 mm in length (Fig. 3B). Germ cells were still visible individually, although some appeared to form clusters. At 3 wpf, the putative gonad was easily distinguishable using transmitted light and was further confirmed using epifluorescent light and histology. The gonad could be precisely dissected from this stage onward. The germ cells had now formed clusters composed of type I and meiotic type II germ cells and an ingrowth of stroma cells (Fig. 3C).

At 6 wpf, ovaries were easily distinguishable in live fish and were positioned between the comparatively darker digestive tract and the swim bladder (Fig. 4A). The ovary consisted predominantly of oocytes in the primary growth stage with both pre-follicle (Ia) and follicle phase (Ib) oocytes present. At 9 wpf, the ovaries had increased in mass and occupied the majority of the body cavity (Fig. 4B). Higher intensity eGFP signal in the ventral portion of the gonad corresponded to primary growth oocytes, whereas oocytes in the cortical alveolus stage (II) and vitellogensis (III) were observed with increased frequency in the dorsal region. In addition, this stage was characterized by the development of the zona radiata and the formation of the theca and granulosa layers. At 12 wpf, the fish had reached sexual maturity.

**Figure 2** Comparison of primordial germ cell visibility at 2 days post fertilization between the (A) See-Thru-Gonad and (B) zf45Tg zebrafish lines. Left column – side view; center column – dorsal view; right column – magnification of primordial germ cells visible at dorsal view. Primordial germ cells in See-Thru-Gonad embryos are seen in better resolution and detail.

**Figure 3** Visualization of the zebrafish germline before sexual differentiation. (A) 4 days post fertilization: lateral and dorsal images allow clear viewing of bright eGFP-expressing cells presumed to be germ cells. (B) 2 weeks post fertilization: the gonad is not visible under transmitted light. Increased magnification and the use of epifluorescent light reveal the proliferating germ cells. (C) 3 weeks post fertilization: the gonad is demarcated under transmitted and epifluorescent light (arrowheads). A comparison of histology and morphology of excised gonad identities eGFP-expressing cells to be type I (Gc1) and meiotic type II germ cells (Gc2). Str, stroma cells.
and the body cavity was filled with vitellogenic and mature (IV) oocytes (Fig. 4C). Under epifluorescent light, vitellogenic oocytes were distinguishable by numerous lipid droplets, while mature oocytes had a comparatively dark appearance.

It was not possible to observe testis under the transmitted light during their development. However, the GFP signal was seen in vivo under epifluorescent light (Fig. 5). At 6 wpf, the testis consisted primarily of clustered spermatogonia and spermatocytes in various stages of development (Fig. 5A). At 9 wpf, large clusters of spermatids were visible (Fig. 5B). The corresponding whole testis images showed clusters of either spermatocytes or spermatids.
surrounded by spermatogonia, which were defined by their comparatively bright GFP signal. At 12 wpf, males were sexually mature (Fig. 5C) and the testes were more noticeable compared with younger fish, and spermatozoa were observed among well-defined clusters of cells in various stages of spermatogenesis.

**Knockdown of miR-92a-3p**

The effect of the VMO treatment was dose-dependent and vertically transmitted. Injection of 0.5 mM VMO at a dose of 3.0 µL directly into ovaries resulted in a significantly lower relative expression of miR-92a-3p in embryos originating from the VMO ovaries, as compared with control embryos treated with a 5-base mismatch control VMO. By comparison, the treatment effect was not significant (P > 0.05) when injecting a 2.0 µL volume and the variation between batches was considerably greater (Fig. 6A).

Arrested development at the 1-cell stage was the primary phenotype associated with the treatment (Fig. 6B). On average, 30.6 ± 13.3% (mean ± s.d.) and 26.4 ± 31.8% of embryos from the 3.0 and 2.0 µL dose treatment groups respectively were arrested at the 1-cell stage, whereas in control groups it was respectively 3.7 ± 2.3% and 1.9 ± 3.6% (Fig. 6C). In both VMO concentrations, the effect of treatment to ovaries, manifested in the developmental arrest frequency, was significant (F_{1.4} = 16.9, P = 0.01 and F_{1.10} = 7.6, P = 0.02, for 3 and 2 µL injection doses respectively). There were no significant effect of treatment on the number of embryos per spawning, and variation in embryos per spawning was large (Fig. 6D). Detailed summary statistics are given in Supplementary Table 2.

**Discussion**

In order to use zebrafish as a model for reproductive biology, we created See-Thru-Gonad, a transparent zebrafish line with a fluorescently labeled germline, visible from 1 dpf onward. Once established and validated, the line was used to test whether the effects of targeted gene knockdown in an ovary can persist in the offspring.

**Genomic validation of See-Thru-Gonad mutations**

Because of the similarity between roy (−/−) and transparent (−/−) zebrafish mutants, and an uncertainty in the background of the mutations in our fish (communication with Zebrafish International Resource Center), we determined the genomic origin of mutations in See-Thru-Gonad. We confirmed that See-Thru-Gonad contains the vasa:vasa-EGFP transgene, while also being homozygous for mutations in the mitfa (nacre) and mpv17 (transparent) genes. While the transcript from mutant mpv17 had been sequenced previously, the researchers were unable to amplify the gene from genomic DNA (Krauss et al. 2013). Here, we demonstrate that the
mutation is the result of an insertion of approximately 3 kb in size located in the second exon of mpv17 (Supplementary Fig. 1).

\textbf{Tg(vasa:vasa-EGFP) has germline-specific expression throughout development}

The See-Thru-Gonad zebrafish line expresses the \textit{Tg(vasa:vasa-eGFP)} specifically in the germline throughout the zebrafish lifecycle, including spermatozoa (Figs 1, 2, 3, 4 and 5). During embryogenesis, and in agreement with previous observations (Krøvel \& Olsen 2002), the expression of the transgene matched the expression pattern of \textit{vasa}, which is specific to the germline (Yoon et al. 1997). Post-hatch, we found the GFP signal to be restricted to the presumptive gonad, which we confirmed both by its excision and through sectioning for histology and comparing with previous studies on zebrafish gonad development (Selman et al. 1993, Maack \& Segner 2003, Neumann et al. 2011).

Two splice variants of \textit{vasa} were identified previously (Krøvel \& Olsen 2004). Pre-zygotically, until approximately 3 wpf, only the shorter variant, \textit{vas-\delta4}, is present in the germline; after sexual differentiation, both longer, \textit{vas-l}, and shorter variants are found in the ovary, while only the shorter variant is present in the testis (Krøvel \& Olsen 2004). Abundance of \textit{vas-l} and \textit{vas-\delta4} was related to the strength of fluorescent signal in zf45Tg fish, with the latter variant producing only a faint signal (Krøvel \& Olsen 2004). In this study, we also experienced that intensity of fluorescent signal in the male germline was weaker; nevertheless, the signal could easily be detected in the See-Thru-Gonad males.

\textbf{See-Thru-Gonad provides an enhanced model for in vivo viewing of the germline}

The See-Thru-Gonad line allows high-resolution imaging of the germline in vivo. During embryogenesis, PGCs actively migrate from their site of specification to the developing gonad, typically arriving within 24 hpf to form clusters on either side of the lateral mesoderm (Yoon et al. 1997, Weidinger et al. 2002). With See-Thru-Gonad, we were able to identify individual PGCs at 2 dpf (Fig. 2A). At this time, several of the embryos examined had individual PGCs that had yet to complete their migration but may not have been detected in zf45Tg fish due to being obscured by pigments.

Post-hatch, observations at 4 dpf and 2 wpf were still able to distinguish individual PGCs in vivo. During this period PGCs are proliferating, and the threshold number of PGCs formed during this period determines the future sex of zebrafish (Siegrfried \& Nüsslein-Volhard 2008, Tzung et al. 2015). In the study by Tzung et al. (2015), PGC counts from excised gonads at 14 dpf were sexually dimorphic, and showed some correlation with the respective sex ratios of their zebrafish families (Tzung et al. 2015). Use of See-Thru-Gonad would allow for observation of PGCs in live individuals, with the ability to track the same individual through sexual maturation.

After sexual differentiation, the See-Thru-Gonad line allowed us to observe the structure of the ovary based on the relative intensity of the eGFP signal. Oocytes in the primary growth stage had noticeably higher eGFP intensity than oocytes in the cortical alveoli and vitellogenic stages, thus making it possible to evaluate the stage of ovary development in vivo (Fig. 4). In males, we were also able to in vivo detect and evaluate oocytes (Fig. 5). Although eGFP intensity was relatively low and required either a high magnification or longer exposure time than in the case of ovaries, See-Thru-Gonad would still be useful for quickly sexing fish, or locating and observing crude morphology of the testis.

These traits provide a significant improvement over the transparent \textit{Tg(bactin:eGFP)}-based line \textit{\beta-ruby}, which does not allow identification of germ cells in fish younger than 2 wpf or in males (Hsiao \& Tsai 2003, Akhter et al. 2016). In addition, the \textit{Tg(vasa:vasa-eGFP)} transgene is germline-specific (Krøvel \& Olsen 2002), while GFP signal in \textit{\beta-ruby} has been detected in the ovary, gills, as well as a small cluster of cells near the genital pore (Akhter et al. 2016).

The uc1Tg and See-Thru-Gonad lines show similar eGFP expression patterns throughout development, but they differ primarily in the initiation of zygotic transcription. While \textit{ziwi:eGFP} transcription begins at 2 dpf, the \textit{vasa:vasa-eGFP} signal is maternal until 3 wpf, with the initiation of meiosis (Krøvel \& Olsen 2004, Leu \& Draper 2010). In our study, we found maternal GFP sufficiently labeled proliferating germ cells before meiosis (Fig. 3). Leu and Draper (2010) demonstrated that stage la oocytes which have yet to enter the diplotene stage of meiotic prophase I do not transcribe \textit{Tg(vasa:vasa-eGFP)} and may not be detected after the maternal signal fades. However, we suggest that See-Thru-Gonad may provide additional flexibility for researchers, since crossing See-Thru-Gonad males with \textit{mitf\textsuperscript{aw2/w2}; mpv17\textsuperscript{b1/b18}} females would create a transparent hybrid allowing for the observation of the onset of meiosis in live fish.

\textbf{Morpholino injections into ovaries result in reduced miR-92a-3p expression in embryos}

In this study we demonstrated that injection of a VMO targeting miRNA miR-92a-3p directly into the zebrafish ovary resulted in significantly decreased abundance of mature miR-92a-3p in 1-cell stage embryos, as well as a significantly increased proportion of embryos with arrested development (Fig. 6). Previously, morpholino-based targeted gene knockdown of the protein-coding gene \textit{incenp} was performed in oocytes of the frog (\textit{Xenopus laevis}) before fertilization, but it required the excision and direct injection of the oocytes before.
transplanting them back into ovulating females (Leblond et al. 2012). To our knowledge, this study is the first demonstration of the vertical transmission of the effect of targeted knockdown of a miRNA, as well as the first time an injection of a VMO to the ovary was observed to have a vertical transmission of the effect to the subsequent embryos.

It is unclear whether the transmission of miR-92a-3p to oocytes occurred after the final maturation, or during vitellogenesis. Because morpholino oligomers are capable of targeting both mature and precursor miRNA, it would be beneficial to explore the composition of these forms of miR-92a transcripts present in oocytes during maturation. We made attempts to collect unfertilized eggs and thus measure the relative expression of mature miR-92a-3p before fertilization. Unfortunately, we were unable to collect sufficient quantities of eggs from injected fish for analysis. Regardless, our findings suggest a novel tool for examining the function of maternal miRNA early in embryonic development.

miR-92a-3p belongs to the conserved miR-17/92 cluster, which in mammals has been linked to regulation of the cell cycle, proliferation and apoptosis (Mogilyansky & Rigoutsos 2013). In mammalian models, one function of miR-92a-3p is to regulate cell proliferation by promoting the cell cycle transition from the G1 to S phase (Manni et al. 2009, Zhou et al. 2015). Another study has shown mammalian miR-92a-3p to regulate insulin expression and maintain glucose homeostasis (Setyowati Karolina et al. 2013). During zebrafish embryogenesis, miR-92a-3p regulates endoderm specification, left/right axis formation as well as control the BMP and nodal signaling pathways through its interaction with their antagonists (Li et al. 2011, Ning et al. 2013). However, the role of miR-92a-3p during ovary maturation and early embryo cleavage has yet to be explored.

Our findings suggest a correlation between miR-92a-3p abundance and the ability of the embryo to initiate cleavage. Reasons for a zebrafish embryo failing to initiate cleavage may include a failure to complete meiosis, a lack of fertilization or a block to the initiation of the first cell cycle (Dosch et al. 2004). Some of the predicted zebrafish miR-92a-3p targets (Vejnar & Zdobnov 2012, Agarwal et al. 2015) are highly expressed zebrafish genes in unfertilized oocytes and during the first cell divisions (Aanes et al. 2011), and have functions related to meiosis and the cell cycle. These include wee1 homolog 2 (wee2), cyclin a1 (ccna1) and eukaryotic translation initiation factor 4E nuclear import factor 1 (eif4enf1). In mammals, Wee2 encodes a conserved oocyte-specific meiosis inhibitor essential to maintaining oocytes in meiotic arrest (Han et al. 2005, Hanna et al. 2010, Oh et al. 2010). In the mouse (Mus musculus), Ccna1 is expressed exclusively in the germ cell lineage and is essential for spermatocyte passage into the first meiotic division (Liu et al. 1998), while in frog, addition of cyclin A to eggs resulted in stable arrest during metaphase of the cell cycle (Roy et al. 1991). Also in mouse, Eif4enf1 is essential for the progression of meiosis, although the mechanism through which it acts is still unclear (Plender et al. 2015). Future studies will attempt to confirm the predicted relationship between zebrafish miR-92a-3p and its predicted targets with possible functions in regulating meiosis and the cell cycle.

**Conclusion**

The See-Thru-Gonad line offers germline-specific signal throughout zebrafish development and enables high-resolution observation of germ cell development in vivo. In addition, the application of See-Thru-Gonad as a resource for reproductive biology was demonstrated by performing targeted gene knockdown of miR-92a-3p in mature oocytes and measuring the treatment effects in embryos after natural spawning. Embryos from treated fish had significantly reduced expression of miR-92a-3p and significantly increased proportion of embryos arrested at the 1-cell stage. For the first time, the effects of targeted gene knockdown using VMOs have been observed to have vertical transmission to offspring.

**Supplementary data**

This is linked to the online version of the paper at http://dx.doi.org/10.1530/REP-16-0328.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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