ENU-Induced Mutagenesis in Grass Carp (Ctenopharyngodon idellus) by Treating Mature Sperm

Xia-Yun Jiang, Cheng-Fei Sun, Quan-Gen Zhang, Shu-Ming Zou*

Key Laboratory of Aquatic Genetic Resources and Utilization, Shanghai Ocean University, Shanghai, China

Abstract

N-ethyl-N-nitosourea (ENU) mutagenesis is a useful approach for genetic improvement of plants, as well as for inducing functional mutants in animal models including mice and zebrafish. In the present study, mature sperm of grass carp (Ctenopharyngodon idellus) were treated with a range of ENU concentrations for 45 min, and then wild-type eggs were fertilized. The results indicated that the proportion of embryos with morphological abnormalities at segmentation stage or dead fry at hatching stage increased with increasing ENU dose up to 10 mM. Choosing a dose that was mutagenic, but provided adequate numbers of viable fry, an F1 population was generated from 1 mM ENU-treated sperm for screening purposes. The ENU-treated F1 population showed large variations in growth during the first year. A few bigger mutants with morphologically normal were generated, as compared to the controls. Analysis of DNA from 15 F1 ENU-treated individuals for mutations in partial coding regions of igf-2a, igf-2b, mstn-1, mstn-2, fst-1 and fst-2 loci revealed that most ENU-treated point mutations were GC to AT or AT to GC substitution, which led to nonsense, nonsynonymous and synonymous mutations. The average mutation rate at the examined loci was 0.41%. These results indicate that ENU treatment of mature sperm can efficiently induce point mutations in grass carp, which is a potentially useful approach for genetic improvement of these fish.

Introduction

Genetic breeding of aquacultural fish species mainly depends on the discovery of natural mutants with high performance, and then improved strains can be produced through genetic selection, hybridization or marker-assisted breeding approaches [1,2,3]. To date, 73 improved breeds of aquacultural species have been produced in China, among which 39 strains were created by selective breeding. All of these improved breeds were developed from existing natural mutants that carry desirable genes or traits [4]. All of these improved breeds were developed from existing natural mutants that carry desirable genes or traits [5,6]. Thus, the acquisition of mutants with high performance is key for achieving breeding goals. However, the rate of natural spontaneous mutations in fish species is generally lower than \(10^{-3}\), meaning that chemical mutagenesis is an efficient way to produce new mutants for future genetic improvement in aquacultural species [7].

Farmed fish are fertilized in vitro and display high reproductive abilities, which allows for chemical mutagenesis of the sperm, egg or embryo at various stages [7,8,9]. N-ethyl-N-nitosourea (ENU) is a chemical mutagen that acts as an alkylating agent, transferring its ethyl group to nucleophilic nitrogen or oxygen sites on deoxyribonucleotides, leading to base mismatch during DNA replication [10,11]. ENU treatment mainly induces single-base substitutions that resemble natural spontaneous mutations [9]. Therefore, the mutation rate induced by ENU at specific loci ranges from \(0.5 \times 10^{-3}\) to \(3.9 \times 10^{-3}\) in mice and zebrafish, which is almost 10 times greater than mutation rates induced using any other means [12,13]. In addition, mutations induced by ENU are unbiased meaning that all genes are mutated at random [14]. Induction of mutants using ENU has become an important method for examining the functional genome of model organisms including fruit flies, zebrafish, medaka, clawed frog and mice [15,16]. Furthermore, though chemical mutagenesis has been used widely in genetic breeding of microorganisms and crops, the application of ENU in fish breeding has been reported only rarely [17,18].

The grass carp, Ctenopharyngodon idellus, is an important herbivorous fish species for freshwater culture, with total annual production of 3.56 million tons in China alone [19]. The brood fish take at least five years to attain sexual maturity, and bodyweight usually exceeds 10 kg. Recently, Chinese investigators had made large effort in understanding genetic variations in natural and cultured populations using DNA markers and in developing genomic tools to facilitate breeding [20,21,22,23,24]. To date, however, breeds of grass carp with improved characteristics have not been produced by conventional mass selection due to its long life cycle. In this present study, we performed in vitro chemical mutagenesis of postmeiotic sperm, rather than in vivo spermatogonial treatment in grass carp, as this former method is easier in fish species with large bodyweights and it gives a high efficiency of mutation induction. After examining embryo development, first year growth and base changes of six growth-related genetic loci in an F1 population, ENU mutagenesis was shown to be efficient for inducing point mutations in the genome.
of the grass carp. In addition, large growth variations existed in the ENU-treated F1 population, which suggests that ENU mutagenesis may be a useful approach for future genetic breeding programs in grass carp.

Materials and Methods

Ethics Statement

This study was approved by the institutional review board or ethics committee of Shanghai Ocean University (Permit Number: 2009011). All experiments were conducted following guidelines approved by the Shanghai Ocean University Committee on the Use and Care of Animals.

ENU working solution

All working solutions were freshly prepared following the methods described by Jin et al. [14]. Briefly, 0.1 g of ENU (Sigma) was dissolved in 0.5 mL of 10 mmol/L sodium acetate to give a 0.1 mol/L stock solution. Before use, the ENU stock solution was dissolved in modified Hanks solution to give 0.5, 1, 5 and 10 mmol/L working solutions. The modified Hanks solution contained 0.4 g KCl, 8 g NaCl, 0.35 g NaHCO3, 0.09 g NaH2PO4·2H2O, 0.1 g MgSO4·7H2O, 0.1 g MgCl2·6H2O, 0.06 g KH2PO4, 0.14 g CaCl2 and 1 g glucose in 1 L sterile deionized water.

Experimental fish

The brood stock of grass carp was maintained at the Qingpu fish breeding facility of the Shanghai Ocean University. Well-developed 6-year-old broodfish were selected for use (four males and four females; mean body weight of 12 kg). The females were injected with 4 μg/kg luteinizing hormone-releasing hormone A2 (LHRH-A2, Ningbo Hormone Company, China), while the males received only half this dose. The broodfish were kept in a circular breeding pond (4 m in diameter) with flowing water stimulation. Meanwhile, the female fish were kept in the breeding pond during the sperm treatment.

ENU treatment, fertilization, hatching and cultivation

Four milliliters of each ENU working solution was mixed with 1 mL grass carp milts in 15-mL screw-cap centrifuge tubes and then incubated at room temperature for 45 min. Sperm activity was examined every 10 min under a light microscope. After ENU treatment, grass carp eggs were manually stripped into a dry bowl and then these were treated immediately with ENU working solutions. Meanwhile, the female fish were kept in the breeding pond during the sperm treatment.

Table 1. Primers used in this study and their related amplification region in grass carp (Ctenopharyngodon idellus).

| Locus | Primers 5‘-3’ | Size (bp) | Exon | Coding region (bp) |
|-------|--------------|-----------|------|-------------------|
| igf-2a | Forward: acaacagaggcttccaagaaa <br> Reverse: tcaacttgggctcaacgtagt | 263 | 4 | 389–651 |
|       |              |           |      |  |  |
| igf-2b | Forward: tgtgaagttaccaataaaga <br> Reverse: tcaacttgggagttctga | 214 | 4 | 393–606 |
| mstn-1 | Forward: atgcagtttacagctgtt <br> Reverse: gcctgtgatgctctgttagt | 396 | 1-396 |  |
| mstn-2 | Forward: caactcatcactacatgtt <br> Reverse: cagtcctttctctcctgg | 369 | 2 | 358–726 |
| fis-1 | Forward: atggccagagtctgtgag <br> Reverse: cctagctgggacgggct | 236 | 5 | 731–966 |
| fis-2 | Forward: agaagccagagtcttgagtg <br> Reverse: aagtgccagggcgggct | 213 | 5 | 738–950 |

Determination of F1 mutation sites

Total genomic DNA was isolated from fish fin clips (0.1 to 0.2 g) using a standard phenol-chloroform procedure detailed by Sambrook et al. [25]. Primers were derived from the published open reading frame sequences of igf-2a, igf-2b, mstn-1, mstn-2, fis-1 and fis-2 (Table 1), which were selected as growth-related genes [26,27].

The PCR reactions (25 μL) contained 10 mmol/L Tris-HCl (pH 8.4), 20 mmol/L KCl, 10 mmol/L (NH4)2SO4, 1.5 mmol/L MgCl2, 0.1 mmol/L of each dNTP, 0.2 μmol/L of each primer, ca. 200 ng gDNA and 2 U pfu Tag DNA polymerase (Applied Biosystems). PCR thermal cycles comprised of one cycle of pre-denaturation (94°C for 5 min), followed by 35 cycles of amplification (94°C for 30 s, 50°C for 30 s, 72°C for 45 s), and a final extension step (72°C for 5 min). PCR products (3 μL) were analyzed by agarose gel electrophoresis, stained with ethidium bromide, and photographed using a Bio-Rad gel image system.

Direct sequencing of the amplified regions of the six selected gene loci was performed in the parents, which can prevent noisy background from single nucleotide polymorphisms. PCR was carried out by forward primers (5’- GGTGTAACCTAGTCGTACTA -3’ for igf-2a, 5’- GGGAGATGTTTCTCCACATG -3’ for igf-2b, 5’- CGTGCGTGGTTGAGGTCTATTG -3’ for mstn-1, 5’- AGGATGAGGAAACAGGATG -3’ for mstn-2, 5’- GGCACCGATGGATTGTTTAC -3’ for fis-1 and 5’- TGCATGCAGAAGGGCAGACTT -3’ for fis-2) and corresponding reverse primers in Table 1. PCR products were recovered from the gel and sequencing were performed on a 48 capillary 3730 DNA Analyzer (Applied Biosystems, Foster City, CA) using aforementioned forward primers following the methods described by Xu et al. [28]. To increase the probability of detecting mosaic mutation sites in the ENU-treated F1 progeny, which interferes with direct sequence analysis, clone sequencing of amplified regions was chosen to detect mutations at each gene locus. PCR products were recovered from the gel and cloned into the pMD19-T (TaKaRa) vector using Escherichia coli DH5α cells. Eight positive clones from each gene fragment were sequenced on a 48 capillary 3730 DNA Analyzer. Mutations were identified using software based on Polyphred version 6.0 Beta [29], which compares each trace to the parental reference sequence and identifies potential
mutations. If a base change was seen in more than two eighths
of the clones for each gene amplicon, it was categorized as a
mutation. The following equation was used to calculate the
mutation rate at each locus:

\[
\text{Mutation rate} \% = \frac{\text{Number of mutated sites}}{\text{Number of fish examined} \times \text{Locus size (bp) excluding primers}}
\]

**Statistical analysis**
Differences between groups were analyzed using one-way
analysis of variance tests, followed by Fisher's post hoc test or
unpaired \(t\)-tests. Significance was accepted at \(p<0.01\).

**Results**

**Effect of ENU treatment on embryo development of grass carp**
A range of ENU concentrations up to 10 mM was used to treat
mature sperm for 45 min. These sperm were used to fertilize wild-
type grass carp eggs, and the resulting embryos were scored for
dominant effects and viability at the hatching stage. Treatment of
mature sperm with 0.5, 1, 5, 10 mM ENU caused the formation of
16.1%, 38.7%, 66.9% and 91.3% abnormal embryos at the
segmentation stage respectively, and these proportions were
significantly (\(P<0.01\)) greater than the 2.8% seen in the control
group (Table 2). Additionally, the survival rates at the hatching
stage were 76.9%, 52.6%, 14.4% and 4.4% for the embryos
created using sperm treated with 0.5, 1, 5, 10 mM ENU
respectively, which were significantly (\(P<0.01\)) lower than the
93.4% survival seen in the control group (Table 2). Thus, the
proportion of morphologically abnormal and dead fry during
embryogenesis increased with increasing ENU concentration.

**Table 2. Effects of different ENU concentrations on morphology of embryos and hatching success after treating mature sperm of grass carp.**

| ENU concentration (mM) | Crossing group | No. of embryos | Abnormality rate at segmentation stage (%) | Survival rate at hatching stage (%) |
|------------------------|----------------|----------------|------------------------------------------|-------------------------------------|
| 0.5                    | R1 x R1        | 674            | 12.4                                     | 81.8                                |
|                        | R2 x R2        | 653            | 14.6                                     | 76.2                                |
|                        | R3 x R3        | 764            | 21.3                                     | 72.8                                |
|                        | Mean           | 764            | 16.1*                                    | 76.9 *                               |
| 1                      | R1 x R1        | 965            | 33.6                                     | 55.6                                |
|                        | R2 x R2        | 654            | 42.3                                     | 52.4                                |
|                        | R3 x R3        | 563            | 40.4                                     | 49.8                                |
|                        | Mean           | 727            | 38.7 b                                    | 52.6 b                               |
| 5                      | R1 x R1        | 567            | 60.6                                     | 19.6                                |
|                        | R2 x R2        | 845            | 68.2                                     | 13.2                                |
|                        | R3 x R3        | 265            | 71.9                                     | 10.4                                |
|                        | Mean           | 559            | 66.9 c                                    | 14.4 c                               |
| 10                     | R1 x R1        | 765            | 89.3                                     | 2.8                                 |
|                        | R2 x R2        | 565            | 88.2                                     | 9.3                                 |
|                        | R3 x R3        | 365            | 96.4                                     | 1.2                                 |
|                        | Mean           | 565            | 91.3 d                                    | 4.4 d                                |
| 0 (control)            | R1 x R1        | 745            | 2.8                                      | 93.6                                |
|                        | R2 x R2        | 435            | 3.4                                      | 92.5                                |
|                        | R3 x R3        | 542            | 2.3                                      | 94.2                                |
|                        | Mean           | 574            | 2.8 *                                    | 93.4 *                               |

Different letters in the same column represent a significant difference between
two groups (\(P<0.01\)).
doi:10.1371/journal.pone.0026475.t002

**Figure 1. Morphological defects of ENU-treated F1 embryos at hatching stage.** (A) Morphologically normal grass carp embryo. (B) Embryo with cardiac displacement. (C) Embryo with short spine and small head. (D) Embryo with short spine, small head, crooked tail and enlarged pericardial cavity. (E) Embryo without head. (F) Embryo with nervous system abnormalities, including a small head and crooked tail. Scale bar = 600 \(\mu\)m.
doi:10.1371/journal.pone.0026475.g001
Growth variation in ENU-treated F1 individuals during the first year

As shown in Table 2, treatment of sperm with 1 mM ENU could produce adequate numbers of viable F1 individuals, which meanwhile displayed substantial dominant mutation effects. Thus, an F1 population was generated for screening purposes using sperm treated with this dose of ENU. The 1 mM ENU-treated F1 populations were derived from three pairs of parents as shown in Table 2. The ENU-treated F1 populations and untreated controls were reared separately in six earth ponds with the same conditions. After eight months of rearing, fish were captured, labeled with PIT, and then bodyweights were determined. Approximately 50% of the 1 mM ENU-treated F1 individuals were morphologically normal (Fig. 2A and 2B), but the others displayed various extents of development retardation and body defects (Figs. 2C, 2D and 2E). As shown in Fig. 3A, the bodyweights of the ENU-treated F1 individuals ranged from 204.5 to 756.6 g, with a mean bodyweight of 437.1 ± 276.2 g (± one standard deviation) (Table 3). The bodyweights of F1 control fish ranged from 304.2 to 576.4 g, with a mean bodyweight of 548.7 ± 42.4 g (Table 3, Fig. 3B). Although the mean bodyweight of the ENU-treated F1 population was only 80% of the control fish, the standard deviation of bodyweight was 6.5-fold greater in the ENU-treated F1 population than in the control population. In the ENU-treated F1 population, more than 85% (478/560) of morphologically abnormal individuals were of lower (<600g) bodyweight, while a portion of 62% (166/274) progenies with bodyweight bigger than 600g were morphologically normal (Fig. 3). These morphologically normal mutants with high growth rates in the ENU-treated F1 populations may be useful in future breeding.

Detection of mutation sites and frequencies in the ENU-treated F1 population

To determine the mutation sites and frequencies in the F1 population derived from the 1 mM ENU-treated sperm, genomic DNA was isolated from 15 ENU-treated F1 fish and five control fish, which were derived from a single pair of parents. Sequences of six gene loci (igf-2a, igf-2b, mstn-1, mstn-2, fst-1 and fst-2) were analyzed. As shown in Table 4, the amplified fragments (excluding the primers) were 224 bp, 174 bp, 356 bp, 329 bp, 196 bp and 173 bp for igf-2a, igf-2b, mstn-1, mstn-2, fst-1 and fst-2 respectively. Among the 15 ENU-treated individuals, there were 15, 10, 18, 21 and 12 point mutations at these loci respectively, while the mutation rates were 0.45%, 0.38%, 0.34%, 0.43%, 0.41% and 0.46% respectively (Table 4). No mutations were identified at these gene loci in the F1 control individuals.

Among the nucleotide substitutions seen at the six selected gene loci in ENU-treated F1 individuals, 52% (46/88) were GC to AT transitions, 35% (31/88) were AT to GC transitions, 9% (8/88) were AT to TA transversions, two was an AT to CG transversion and one was a GC to TA transversion (data not shown). These substitutions led to nonsynonymous changes in approximately 66% (50/88) of cases, of which approximately 64% (31/50) were missense changes, while the remainder gave nonsense mutations. As shown in Fig. 4, an individual with PIT 690,000,116,601,909 (see Fig. 2B) had a C to T point mutation at nucleotide 205 within the 190 bp to 252 bp region of mstn1, which resulted in the Gln at position 69 being substituted with a stop codon (Fig. 4A and 4B). The individual with PIT 690,000,116,601,814 (see Fig. 2A) had an A to G mutation at nucleotide 239, which resulted in a Gln to Arg substitution at position 80 (Figs. 4A and 4C). Additionally, within the coding region of 652 bp to 726 bp of mstn2 an individual with PIT 690,020,042,302,468 (see Fig. 2E) showed a G to A mutation at nucleotide 658, which resulted in a Val to Ile substitution at position 220 (Fig. 5A and 5B). A further individual with PIT 690,020,042,302,463 (see Fig. 2D) showed a G to A change at nucleotide 717, which resulted in a synonymous substitution (Figs. 5A and 5C).

Discussion

Compared to the low natural spontaneous mutation frequency in fish, chemical mutagenesis can be useful for increasing genetic mutations. Highly efficient mutagens can induce profound changes in genetic material and produce mutants with desirable traits. Studies with ENU in the cyprinidae model species zebrafish have obtained mutations in hundreds of genes that are required for embryonic viability [30,31]. ENU-induced mutants for targeted genes have also been reported for medaka [32,33]. Although ENU mutagenesis is a potential method for genetic breeding in commercial fish species, the successful generation of improved
carp strains by ENU mutagenesis combined with gynogenesis has only been reported by Kirpichnikov [18]. Recently, Han et al. [34] reported ENU mutagenesis studies in the gibel carp (Carassius gibelio Bloch) by in vivo spermatogonial treatment. In this present study, application of ENU mutagenesis in the grass carp by treating mature spermatozoa was explored. The 1 mM ENU-treated F1 population of the grass carp showed greater variations in bodyweight during the first year of growth compared with the control group. Although the mean bodyweight of the ENU-treated F1 population decreased to 80% of the controls, the standard deviation of bodyweight in the ENU-treated group was 6.5-fold greater than the controls. A few morphologically normal, but bigger individuals in the ENU-mutated F1 population may provide a useful resource for further functional gene identification studies and genetic improving programs of grass carp.

ENU mutagen treatments can influence the fertilization abilities of sperm [15]. By discarding embryos that displayed defects prior to neurulation stages, it is possible to efficiently screen for later phenotypes in patterning, organogenesis and differentiation. In the present study, the proportion of morphologically abnormal or dead hatching stage embryos increased with increasing dose of ENU. ENU mutagenesis produced various dominant mutations.

Table 3. Growth of ENU-treated F1 fish after eight months of rearing.

| Crossing group | ENU concentration (mM) | No. of F1 fish | Body weight range (g) | Body weight (g) |
|----------------|------------------------|---------------|-----------------------|-----------------|
| g1 × c1        | 1                      | 304           | 218.3–756.6           | 431.2±270.1     |
|                | 0 (control)            | 102           | 511.7–576.4           | 555.8±44.7      |
| g2 × c2        | 1                      | 322           | 212.1–744.2           | 452.9±292.4     |
|                | 0 (control)            | 128           | 504.2–565.5           | 538.5±40.3      |
| g3 × c3        | 1                      | 418           | 204.5–750.6           | 427.2±266.1     |
|                | 0 (control)            | 189           | 519.3–559.8           | 551.8±42.2      |
| Mean           | 1                      | —             | —                     | 437.1±276.2     |
|                | 0 (control)            | —             | —                     | 548.7±42.4      |

Different letters in the same column represent a significant difference between two groups (P<0.01).
and/or synthetic lethal mutations, in which genetic material has been modified. The dominant phenotypes caused by ENU treatment in grass carp led to the disruption of specific embryonic development processes, such as notochord patterning and nervous system and internal organ development. Nevertheless, both forward and reverse genetic screens will be required to confirm whether the mutations are responsible for the dominant phenotypes observed in the ENU-treated progeny [35]. Once a

---

| Locus  | Size (bp) excluding primers | Group                        | No. of fish examined | No. of clones sequenced/fish | No. of mutated sites | Mutation rate (%) |
|--------|-----------------------------|------------------------------|----------------------|-------------------------------|----------------------|-------------------|
| igf-2a | 224                         | ENU-treated F1              | 15                   | 8                             | 15                   | 0.45              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| igf-2b | 174                         | ENU-treated F1              | 15                   | 8                             | 10                   | 0.38              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| mstn-1 | 356                         | ENU-treated F1              | 15                   | 8                             | 18                   | 0.34              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| mstn-2 | 329                         | ENU-treated F1              | 15                   | 8                             | 21                   | 0.43              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| fst-1  | 196                         | ENU-treated F1              | 15                   | 8                             | 12                   | 0.41              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| fst-2  | 173                         | ENU-treated F1              | 15                   | 8                             | 12                   | 0.46              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |
| Average| 242                         | ENU-treated F1              | 15                   | 8                             | 15                   | 0.41              |
|        |                              | F1 control                  | 5                    | 8                             | 0                    | 0                 |

Mutation rate (%) = Number of mutated sites/Number of fish examined x Locus size (bp) excluding primers.

doi:10.1371/journal.pone.0026475.t004

---

Figure 4. Mutation sites at the mstn1 locus. (A) Partial mstn1 sequence alignment of five ENU-treated F1 juvenile fish with sequences from their parents. The numbers by the sequences correspond to the passive integrated transponder tags as shown in Figure 2. (B) Location of the MSTN1\textsuperscript{Q80R} mutation and sequencing that revealed a C to T point mutation. (C) Location of the MSTN1\textsuperscript{Q80R} mutation and sequencing that revealed an A to G point mutation. Dam and sire denote to female and male, respectively.

doi:10.1371/journal.pone.0026475.g004
In the present study, mutations were induced in vitro by ENU treatment of postmeiotic sperm. The initial mutation, usually occurred with an ethylation of a base on one DNA strand, leads to mosaic offspring in the next generation that interferes with direct sequence analysis in the F1 generation. To detect mutations in the partial coding regions of *igf-2a, igf-2b, mstn-1, mstn-2, fst-1* and *fst-2*, clone sequencing of amplified regions was performed. The results showed that these loci displayed mainly GC to AT or AT to GC substitutions, resulting in nonsynonymous nonsense or missense mutations (66%), as well as synonymous mutations. The induced point mutations in the grass carp were similar to those seen in mammalian cells [9,36], where most ENU-induced mutations are GC to AT transitions, and to a lesser extent AT to GC substitutions, resulting in nonsynonymous nonsense or missense mutations (66%), as well as synonymous mutations. The induced point mutations in the grass carp were similar to those seen in mammalian cells [9,36], where most ENU-induced mutations are GC to AT transitions, and to a lesser extent AT to GC transitions, although all types of transitions and transversions have been documented after exposure to ENU. Interestingly, neither deletions nor translocations were identified in this present study, however such mutations have been induced in zebrafish by postmeiotic ENU treatment of male germ cells [35]. It is possible that multigene deletions are induced by the protocol used in this present study, and these may result in a higher frequency of early lethal phenotypes, which were discarded. On the other hand, it has also been proposed that subtle differences in mutagenesis conditions may result in significant differences in the kinds of lesions produced [15].

Mutation rates caused by ENU range from $0.5 \times 10^{-3}$ to $3.9 \times 10^{-3}$ at specific loci when *in vivo* spermatogonial treatment has been used in mice and zebrafish [12,13]. Moreover, ENU mutagenesis in medaka after *in vivo* spermatogonial treatment induces mutation rates of $1 \times 10^{-3}$ to $1.9 \times 10^{-3}$ [8,33]. Genetic and molecular tests have shown that postmeiotic ENU treatment can induce point mutations [37], but the range of mutations induced has not been analyzed extensively. In this present study, the average point mutation rate was $4.1 \times 10^{-3}$ at six selected gene loci in grass carp after *in vitro* chemical mutagenesis of postmeiotic sperm. These results indicate that treatment of postmeiotic gametes with ENU induces point mutations at a higher rate than premeiotic regimens, suggesting that postmeiotic mutagenesis protocols could be useful in genetic screening strategies. Postmeiotic mutagenesis has been reported to produce a 10-fold increase in the frequency of induced mutations in specific-locus tests in zebrafish [35].

In summary, the ENU-treatment of mature sperm with different doses can markedly generate dominant effects on embryo development of grass carp. The ENU-treated F1 populations demonstrated large variations in bodyweight during the first year. Some bigger mutants with morphologically normal were produced in the ENU-treated F1 progeny, which may be useful for genetic breeding in future. Our further sequence data showed that the postmeiotic ENU treatment can efficiently induce point mutations. Most of these point mutations were GC to AT or AT to GC substitutions that led to nonsense, nonsynonymous and synonymous mutations. The classical three-generation of breeding or two-generation gynogenetic screen may be used to confirm mutants that carry desirable genes or traits for breeding in future.

**Author Contributions**

Conceived and designed the experiments: XYJ CFS SMZ. Performed the experiments: XYJ CFS QGZ SMZ. Analyzed the data: XYJ CFS SMZ. Contributed reagents/materials/analysis tools: XYJ CFS QGZ SMZ. Wrote the paper: XYJ CFS SMZ.

Figure 5. Mutation sites at the *mstn2* locus. (A) Partial *mstn2* sequence alignment of five ENU-treated F1 juvenile fish with sequences from their parents. The numbers by the sequences correspond to the passive integrated transponder tags as shown in Figure 2. (B) Location of the MSTN2V220I mutation and sequencing that revealed a G to A point mutation. (C) Location of the MSTN2E239E mutation and sequencing that revealed a G to A point mutation. Dam and sire denote to female and male, respectively.

doi:10.1371/journal.pone.0026475.g005
References

1. Khoo SK, Ozaki A, Nakamura F, Arakawa T, Nickolov R, et al. (2004) Identification of a novel chromosomal region associated with infectious hematopoietic necrosis (IHNV) resistance in rainbow trout Oncorhynchus mykiss. Fish Pathol 39: 95–101.

2. Ozaki A, Sakamoto T, Khoo S, Nakamura K, Coimbra MRM, et al. (2001) Quantitative trait loci (QTLs) associated with resistance/susceptibility to infectious pancreatic necrosis virus (IPNV) in rainbow trout (Oncorhynchus mykiss). Mol Genet Gen I 365: 23–31.

3. Ozaki A, Okamoto H, Yamada T, Matuyama T, Sakai T, et al. (2010) Linkage analysis of resistance to Steinernecus iniae infection in Japanese flounder Paralichthys olivaceus. Aquaculture 308: 562–567.

4. Yu X (2011) Bulletin of Ministry of Agriculture: aquacultural new breeds. Mod Fish Inform 26: 29.

5. Fuji K, Kobayashi K, Hasegawa O, Coimbra MRM, Sakamoto T, et al. (2006) Identification of a single major genetic locus controlling the resistance to lymphosarcoma disease in Japanese flounder Paralichthys olivaceus. Aquaculture 254: 203–210.

6. Fuji K, Hasegawa O, Honda K, Kamadsaka K, Sakamoto T, et al. (2007) Marker-assisted breeding of a lymphosarcoma disease-resistant Japanese flounder Paralichthys olivaceus. Aquaculture 272: 291–295.

7. Liu S (1988) A review of chemical mutagenesis of fish. J Fish China 12: 81–86.

8. Shima A, Shimada A (1991) Development of a possible nonmammalian test system for radiation-induced germ-cell mutagenesis using a fish, the Japanese medaka Oryzias latipes. Proc Natl Acad Sci U S A 88: 2545–2549.

9. Knapik, EW (2000) ENU mutagenesis in zebrafish—from genes to complex diseases. Manum Genome 11: 519–31.

10. Mullins MC, Hammerschmidt M, Hafler P, Niueslein-Volhard C (1994) Large-scale mutagenesis in the zebrafish: in search of genes controlling development in a vertebrate. Curr Biol 4: 109–202.

11. van Eden FJ, Granato M, Odenthal J, Hafler P (1999) Developmental mutant screens in the zebrafish. Meth Cell Biol 60: 21–41.

12. Hitooutcharit S, Carpenter DD, Russell WL (1985) Dose-repition increases the mutagenic effectiveness of N-ethyl-N-nitrosourea in mouse spermatogonia. Proc Natl Acad Sci U S A 82: 6619–6621.

13. Driever W, Solnica-Krezel L, Schier AF, Talbot WS (2000) Analysis of chromosomal rearrangements induced by postmeiotic mutagenesis with ethylnitrosourea. Proc Natl Acad Sci U S A 97: 9061–9066.

14. Jin P, Tian T, Sun Z, Meng A (2004) Generation of mutants with developmental defects in zebrafish by ENU mutagenesis. Chinese Sci Bull 49: 2134–2138.

15. Goda T, Aiba-Daya A, Carruthers S, Clark MD, Stemple DL, et al. (2006) Genetic screens for mutations affecting embryogenesis in zebrafish. Development 132: 47–46.

16. Jin P, Tian T, Sun Z, Meng A (2004) Generation of mutants with developmental defects in zebrafish by ENU mutagenesis. Chinese Sci Bull 49: 2134–2138.

17. Lou Y (1999) Fish breeding. 1st edn Beijing: China Agriculture Press. pp 136–137.

18. Kirpichnikov VS (1981) Genetic bases of fish selection. Berlin: Springer-Verlag. pp 302–303.

19. FBMA (2008) Chinese Fisheries Year Book. Beijing: China Agriculture Press. pp 8–10.

20. Boney SE, Shelton WL, Yang SL, Wilken LO (1984) Sex reversal and breeding of grass carp. Trans Am Fish Soc 113: 348–353.

21. Liu F, Xia JH, Bai ZY, Fu JJ, Li JL, et al. (2009) High genetic diversity and substantial population differentiation in grass carp (Ctenopharyngodon idella) revealed by microsatellite analysis. Aquaculture 297: 51–56.

22. Xia JH, Liu F, Zhu ZY, Fu JJ, Fung JB, et al. (2010) A consensus linkage map of the grass carp (Ctenopharyngodon idella) based on microsatellites and SNPs. BMC Genomics 11: 135.

23. Zhang H, Liu SJ, Zhang C, Tao M, Peng LY, et al. (2011) Induced gynogenesis in grass carp (Ctenopharyngodon idella) using irradiated sperm of allotetraploid hybrids. Mar Biotechn 13: 1017–1026.

24. Zhao JL, Cao Y, Li SF, Li JL, Deng YF, et al. (2011) Population genetic structure and evolutionary history of grass carp Ctenopharyngodon idella in the Yangtze River, China. Environ Biol Fish 90: 85–93.

25. Sambrook J, Fritsch EF, Maniatis T (1989) Molecular cloning: A laboratory manual. 2nd ed New York: Cold Spring Harbor.

26. Zhou S, Kamei H, Mosi Z, Duan C (2009) Zebrafish insulin-like growth factor genes: Gene duplication, structural and expression conservation and divergence. PLoS ONE 4: e7026.

27. Yuan X, Jiang X, Pu J, Li Z, Zou S (2011) Functional conservation and divergence of duplicated IGF-2 genes in grass carp (Ctenopharyngodon idella). Gene 470: 46–52.

28. Xi ZY, Bao QY, Niu YX (2002) Factors that influence direct sequencing of PCR products. Hervedia (Beijing) 24: 548–550.

29. Nickerson DA, Tohe VO, Taylor SL (1997) PolyPhred: automating the detection and genotyping of single nucleotide substitutions using fluorescence based ressequencing. Nucleic Acids Res 25: 2745–2751.

30. Li M, Cao Y, Zhao ZN, Lin S, Meng AM (2001) Characterization and expression pattern of pouII1, a novel class POU gene in zebrafish. Chinese Sci Bull 46: 1523–1527.

31. Amsterdam S, Hopkins N (2006) Mutagenesis strategies in zebrafish for identifying genes involved in development and disease. Trends Genet 22: 473–476.

32. Ishikawa Y (2000) Medaka fish as a model system for vertebrate developmental genetics. Bioessays 22: 487–495.

33. Sambrook J, Fritsch EF, Maniatis T (1989) Molecular cloning: A laboratory manual. 2nd ed New York: Cold Spring Harbor.

34. Han Q, Chang Y, Liang L, Gao G, Chi B, et al. (2010) Genetic diversity of F1 individuals of ENU-mutated gibel carp, Carassius auratus gibelio based resequencing. Nucleic Acids Res 25: 6619–6621.

35. Imai Y, Feldman B, Schier AF, Talbot WS (2000) Analysis of chromosomal rearrangements induced by postmeiotic mutagenesis with ethylnitrosourea in zebrafish. Genetics 155: 261–272.

36. ShibaS T, Morimoto K (1993) A review of the genotoxicity of 1-ethyl-1-nitrosourea. Mutat Res 297: 5–38.

37. Riley RB, Grunwald DJ (1995) Efficient induction of point mutations allowing recovery of specific locus mutations in zebrafish. Proc Natl Acad Sci U S A 92: 5997–6001.