Transfer of Green Fluorescent Protein (GFP) Gene to Betta splendens Embryos by Transfection and Electroporation Methods

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ABSTRACT

Transfection and electroporation method shave a high possibility to apply towards transgenic production of small eggs size fish species. This study aimed to examine the potential of transfection and electroporation methods to use for transferring a foreign gene into betta fish (Betta splendens) embryos using green fluorescent protein (GFP) gene as a model. Fish were spawned naturally in the ratio of male: female was 1:1, then a total of 200 eggs were taken for each treatment. Transfection was performed for 30 minutes (room temperature of about 25 °C) at two-cell stage of embryos using transfast reagent. Transfection reaction consisted of 0.75 µL transfast reagent, 0.25 µL GFP expression vector (DNA concentration: 50 µg/µL) and 99 µL NaCl solution (concentration: 0.95%). Electroporation was performed using 125 volt cm-1, 3 times pulse frequency at one second interval and pulse length of 7 micro seconds. A volume of 800 µL GFP expression vector solution (DNA concentration: 50 µg/ µL) in PBS was used for electroporation. The successful of foreign gene transfer was determined by PCR method with GFP specific primers. The results showed that hatching rate of eggs in transfection treatment was 67.08%, while the electroporation was 72.09%. Survival of larvae in transfection treatment was 73.00%, while the electroporation was 75.00%. The results of PCR analysis showed that transfection method allowed 65% of the survived fish carrying GFP gene, whereas the electroporation method was 70%. Thus, foreign gene transfer in betta fish can be conducted using the transfection and electroporation methods.

Keywords: electroporation, embryo, gene transfer, transfection, Betta splendens

1. Introduction

Betta fish (Betta splendens) is a freshwater ornamental fish widely known in Indonesia. Betta fish is popular in Indonesia as fighting fish and ornamental fish. Ornamental betta fish is commonly preferred by people for its beautiful color in the body and fins. Betta fish strain has very much variation based on color and shape of fin.

Beautiful color in betta fish can be obtained through selection, but this method takes quite a long time (5-7 generation). Selection method is generally performed by betta fish culturists, thus various types of betta fish are widely available. A more modern method yet rarely applied in ornamental fish is transgenesis (gene transfer). Application of transgenesis is expected to improve desirable characteristics for aquaculture such as growth rate increase, muscle quality improvement, as well as increase in fish resistance to extreme environment and diseases (Parenrengi et al., 2011). In ornamental fish the character improvement is intended for color and shape.

Transgenesis is applied to introduce new genetic characteristics or over-expression to individual through the introduction of foreign gene connected to the host genome, which is expected to be inherited to the offspring through gonad cell (Yoshizaki 2001). The main principle of this technic is to insert foreign DNA to the nucleus of a target cell and combine it to the host genome (Hidayani et al., 2011).
Several transgenesis methods include microinjection, electroporation, transfer through sperm, biolistic, the use of viral vector, lipofection, and transfection (Harrison et al., 1998; Beaumont & Hoare 2003).

Transfection is a method of gene transfer based on the use of lipid as an agent to transport foreign gene across cell membrane (Yamano et al., 2011). The advantages of gene transfer through transfection method include: easy to apply, high implementation success, and not toxic to embryo. Transfection has lower risk of physical damage to target organism, thus hatching rate will increase (Calderon, 2004).

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Electroporation is a physical process in which electrical field at certain voltage is applied to cell, causing temporary pores on cell membrane. Eventually, cell will be permeable and it is possible for cell to absorb liquid from its surrounding. Electroporation is considered to be the most effective, efficient, and easiest method in gene transfer (Tsai et al., 1997). In fact, this method is able to be applied to many fish eggs simultaneously in a short period of time (Cheng et al., 2002). As a preliminary study in order to modify important characters in betta fish, this research was performed to examine gene transfer success on betta fish embryo using green fluorescent protein (GFP) gene as reporter. This is important, because one of the considerations in transgenesis technology is the selection of the promoter; DNA sequences located in an upstream section genes (Hackett, 1993; Glick & Pasternak, 2003) that will set the place, time and the expression level of a gene.

The aim of this study was to compare the success of transfection and electroporation methods, and to determine the DNA concentration in GFP gene transfer by electroporation in betta fish embryo (Betta splendens).

2. Materials and Method

The design used in this study was Completely Randomized Design with six treatments and three replications of each. Experimental design of GFP gene transfer in betta fish embryo is presented in Table 1. The use of DNA concentrations based on previous research results on comet fish (Hadie et al., 2010).

Betta fish spawning and egg collection
Ornamental betta fish broodstock were obtained from Research Institute for Ornamental Fish Culture (BRBIH) Depok, West Java. Betta fish of plakat strain were used in this study. Moreover, the broodstocks used were at 5 months of age, healthy and had no morphological defect. The broodstocks selected were measured for their weight and length. There were six male and 6 female broodstocks used in this research. Male broodstocks were further transferred to spawning containers in the form of plastic basin and spawning substrate of styrofoam previously cut into 7 cm x 5 cm of size was added. Later, matching process was performed, namely by placing the female broodstock in a plastic glass put inside the spawning containers (Figure 1).

Table 1. Experimental design of GFP transfer gene in betta fish embryo

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>Transfection with the addition of DNA and transfast</td>
</tr>
<tr>
<td>TK</td>
<td>Transfection without DNA and transfast (control)</td>
</tr>
<tr>
<td>E30</td>
<td>Electroporation with the addition of PBS and DNA (30 µg µL⁻¹)</td>
</tr>
<tr>
<td>E50</td>
<td>Electroporation with the addition of PBS and DNA (50 µg µL⁻¹)</td>
</tr>
<tr>
<td>EK+</td>
<td>Electroporation with the addition of PBS and without DNA</td>
</tr>
<tr>
<td>EK-</td>
<td>Without PBS and DNA</td>
</tr>
</tbody>
</table>

Description: PBS (Phosphate Buffer Saline)
Matching activity was done for 1 day. During the process, male fish built bubblenest. After that, female and male fish were placed together in spawning container. Betta fish spawning lasted for 3-4 hours by natural spawning method.

After spawning process, eggs in the bubblenest were immediately collected and counted. Eggs were placed on petri dish and manually counted using plastic pipette and hand-counter. Number of betta fish eggs used in this study amounted to 200 eggs for each replication of treatment, thus total number of egg used reached 3,600 eggs.

Transfection

Transfection was performed during embryo phase of 2-64 cells. Transfection solution used was a mix of plasmid DNA (concentration of 50 µg mL⁻¹) and transfast in 0.9% NaCl media to reach final volume of 100 µL media. In transfection treatment (T), 0.37 µL of plasmid DNA, 0.75 µL of transfast, and 98.89 µL of NaCl were used. Moreover, in control treatment of transfection (TK), there was no addition of plasmid DNA and 0.75 µL of transfast, thus NaCl used was 100 µL. The solution was further added to the embryo until it was completely submerged, and later the embryo was incubated at room temperature (approximately 25 °C) for 30 minutes. This program referred to the transfection method performed on wild betta fish (Prasetio et al., 2013).

Electroporation

Electroporation was done by mixing plasmid DNA and phosphate buffer saline (PBS) to reach final volume of 800 µL. Treatment of E30 plasmid DNA (concentration of 30 µg µL⁻¹) used 13 µL and 787 µL of PBS, treatment E50 plasmid DNA (concentration of 50 µg µL⁻¹) used 22 µL and 778 µL of PBS, treatment EK+ used 800 µL of PBS without the addition of plasmid DNA, and treatment EK- was without the addition of plasmid DNA and PBS. Sample and the mix were put into cuvette which was previously immersed in 95% ethanol solution and dried. Later, sample was placed in the cuvette. Electroporation was done using Gene PulsarII, and electroporation program was set at voltage of 125 V cm⁻¹, pulse frequency of 3 times, pulse length of 7 milliseconds, and interval of 1 second. This program referred to electroporation method performed in pangasius (Dewi et al., 2012) and comet fish (Hadie et al. 2010).

Hatching rate, sampling of weight and length

After treatments were performed, the number of fertilized eggs were counted. This activity was conducted 1-2 hours after spawning process. Fertilized eggs were transparent, while unfertilized eggs had milk white color. Later, eggs were placed in styrofoam bowl with volume of 1 L and equipped with tea filter. Hatching container of betta fish eggs can be seen in Figure 2 below.
Eggs hatched in one to two days later. Hatching rate (HR) was calculated using the formula of Effendie (1997):

\[
HR(\%) = \frac{\text{number of hatched egg}}{\text{number of fertilized egg}} \times 100
\]

Seven days after hatching, larvae were transferred to styrofoam at a size of 40 cm x 30 cm x 25 cm. Measurement of fish weight and length was performed once every two weeks. Total number of sample was 30 fish of each treatment or 10 fish of each replication.

Formula of absolute growth in length was calculated according to the formula used by Effendie (1997):

\[
AGL = L_t - L_0
\]

**Description:**

- **AGL**: absolute growth in length
- **$L_t$**: average fish length at day-$t$
- **$L_0$**: average fish length at day-0 (beginning of maintenance)

Formula of growth rate in weight was measured using the formula applied by Effendie (1997):

\[
GR = \left(\frac{\ln W_t - \ln W_0}{t}\right) \times 100
\]

**Description:**

- **GR**: growth rate in weight
- **$W_t$**: average fish weight at day-$t$
- **$W_0$**: average fish weight at day-0 (beginning of maintenance)
- **$t$**: time of maintenance

**DNA extraction and detection of GFP gene**

DNA extraction was done through pool sample method with total number of sample amounted to 30 betta fish larvae. DNA was extracted using GeneJET Genomic DNA Purification Kit (Thermo Scientific) followed the method in the manual. Later, measurement of DNA concentration was conducted using Genequant.

Confirmation of GFP gene transfer success into cell was done using PCR method with primers of GFP-F 5'-GGTCGAGCTGGACGG- 3' and GFP-R 5'-ACGAACTCCAGCAGG- 3’ (Rajamuddin, 2010). PCR process was run at pre-denaturation temperature of 94 °C for 3 minutes; 35 cycles for denaturation of 94 °C for 30 seconds, annealing of 62 °C for 30 seconds, extension of 72 °C for 1 minute; and final extension of 72 °C for 3 minutes (Rajamuddin 2010). PCR was performed using PCR machine (Biometra-Jerman).

PCR product was separated through electrophoresis with 1.5% agarose gel. Agarose gel was put into electrophoresis container filled with TBE 1x. PCR product was further placed into the wells of agarose gel using micropipette, and DNA marker was inserted as the marker for fragment size of DNA sample. Electrophoresis was performed at voltage of 100 V for 30 minutes. DNA was visualized using Gel Doc UV Transilluminator. Product of GFP gene amplification was at a size of 600 bp (Rajamuddin, 2010).
As an internal control for gene transfer success, PCR method with primers of \( \beta \)-actin-F 3'-TTGGTCTGTGC-AGGACAA-3' and \( \beta \)-actin-R 3'-AAGGATCCACTGTAAGAAAGGGAA-3' (Higashijima et al., 1997). PCR process was run at pre-denaturation temperature of 94 °C for 3 minutes, 30 cycles for denaturation of 94 °C for 30 seconds, annealing of 58 °C for 30 seconds, extension of 72 °C for 30 seconds, and final extension of 72 °C for 3 minutes. PCR was done using PCR machine (Biometra-Gy). Product of \( \beta \)-actin gene amplification was at a size around 300 bp (Higashijima et al., 1997). The use of \( \beta \) actin as an internal control which is house keeping means that the promoter can be active at any time if necessary.

**Test parameter and data analysis**

Hatching rate, survival rate, absolute growth in length, and growth rate in weight were analyzed using One-way ANOVA and followed with post hoc test of Duncan \( (\alpha=0.05) \) using SPSS Statistic 16.0 (IBM, New York, USA).

### 3. Results and Discussion

Analysis result of hatching rate (HR) is presented in Figure 3. Based on the result, there was no significant difference between treatments \( (P>0.05) \). Average hatching rate of each treatment was 67.08%; 68.75%; 73.43%; 70.75%; 72.89%; and 73.56%, respectively. Similar finding was also obtained on average hatching rate of wild betta fish *Betta imbellis* with range of 70%-80% in research using transfection method conducted by Prasetio et al., (2013). Based on the result of One-Way ANOVA on hatching rate, significant value or \( P \)-value of 0.152 \( (P>0.05) \) was obtained, thus it is concluded that there was no significant difference between hatching rates of each treatment.

This shows that the methods performed both do not significant the hatching rate. The same thing was reported by Tsai et al., (2000) that the hatching rate between the fertilized eggs and electroporation sperm and control sperm is relatively the same. Tsai et al., (2000) also mentions that the difference is only 4% greater in control sperm.

**Description:** The same superscript letter on the same test parameter showed no significant different result based on post-hoc test of Duncan \( (p<0.05) \)

**Figure 3.** Hatching rate of betta fish eggs based on treatment of transfection with the addition of DNA and transfast (T), transfection without DNA and transfast (TK), electroporation with the addition of PBS and DNA at concentration of 30 µg µL\(^{-1} \) (E30), electroporation with the addition of PBS and DNA at concentration of 50µg µL\(^{-1} \) (E50), electroporation with the addition of PBS and without DNA (EK+), and electroporation without PBS and DNA (EK-)
Relatively similar to hatching rate was also obtained for larval survival rate. There was no significant difference between the survival value of betta fish larvae at the age of 5 days in both treatments. Result of survival analysis is presented in Figure 4. Result of survival analysis showed that there was no significant different between treatments (P>0.05). Average survival rate of each treatment was 73.50%; 77.14%; 72.33%; 76.67%; 73.30%; and 69.68%, respectively. Thus the method of electroporation and transfection carried out with both programs does not affect the survival of the larvae produced.

Figure 4. Survival rate of betta fish based on treatment of transfection with the addition of DNA and transfast (T), transfection without DNA and transfast (TK), electroporation with the addition of PBS and DNA at concentration of 30 µg µL⁻¹ (E30), electroporation with the addition of PBS and DNA at concentration of 50 µg µL⁻¹ (E50), electroporation with the addition of PBS and without DNA (EK+), and electroporation without PBS and DNA (EK-)

Detection result of GFP and β-actin in the seed carried the GFP gene and in treatment E50, about 66.67% of seed carried the GFP gene, which indicated that gene transfer through electroporation method can be applied in the embryo of betta fish Betta splendens. Gene transfer using electroporation method in betta fish has not been scientifically published, yet electroporation with GFP inserted to the eggs through spermatozoa in comet fish can be detected (Hadie et al., 2010) as well as electroporation with GFP gene for transplantation of testicular cell of tilapia (Barades 2014) which showed that the percentage of fluorescent cell ranged of 48.00-55.33%.
Table 2. Detection of GFP and β-actin on transgenic betta fish based on treatment of transfection with the addition of DNA and transfast (T), transfection without DNA and transfast (TK), electroporation with the addition of PBS and DNA at concentration of 30 µg µL⁻¹ (E30), electroporation with the addition of PBS and DNA at concentration of 50 µg µL⁻¹ (E50), electroporation with the addition of PBS and without DNA (EK+), and electroporation without PBS and DNA (EK-)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Detection of GFP (%)</th>
<th>Detection of β-actin (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>60.00±0.00</td>
<td>100.00±0.00</td>
</tr>
<tr>
<td>TK</td>
<td>0.00±0.00</td>
<td>100.00±0.00</td>
</tr>
<tr>
<td>E30</td>
<td>80.00±20.00</td>
<td>100.00±0.00</td>
</tr>
<tr>
<td>E50</td>
<td>66.67±11.55</td>
<td>100.00±0.00</td>
</tr>
<tr>
<td>EK+</td>
<td>0.00±0.00</td>
<td>100.00±0.00</td>
</tr>
<tr>
<td>EK-</td>
<td>0.00±0.00</td>
<td>100.00±0.00</td>
</tr>
</tbody>
</table>

Description: The same superscript letter on the same test parameter showed no significant different result based on post-hoc test of Duncan (p<0.05)

Figure 5. Electrophoregram of some samples of PCR product using specific primers of GFP (A) and β-actin (B) with DNA template extracted from caudal fin of betta fish. Marker of DNA fragments size (M), electroporation with the addition of PBS and DNA at concentration of 30 µg µL⁻¹ (E30), electroporation with the addition of PBS and DNA at concentration of 50 µg µL⁻¹ (E50), electroporation with the addition of PBS and without DNA (EK+), electroporation without PBS and DNA (EK-), treatment of transfection with the addition of DNA and transfast (T), transfection without DNA and transfast (TK), and control of PCR (N).

Electrophoregram of some samples of PCR-GFP product is shown in Figure 5. Six samples were not detected in transfection treatment, while all samples of β-actin were able to be detected. In treatment E30, 3 samples of DNA-GFP were not detected and all samples of β-actin could be detected, while in treatment E50, there were 5 samples of DNA-GFP that could not be detected yet all samples of β-actin were detected.

Analysis of GFP detection of pool sample embryo betta fish is positive carrying genes. This suggests that the universal β-actin promoter used is able to control GFP expression of betta fish. GFP genes transferred in addition to being able to inserted in the body of betta fish are also able to be detected properly. According to Sarmasik (2003), if the transgenic construction carries a functional promoter, a number of transgenic individuals can be expected to express the transgene. The β-actin gene in fish is housekeeping gene (can be active whenever needed) and the regulator area is widely used in transgenic fish research.

As a supporting parameter in this study is the growth of larvae. Analysis result of absolute growth in length is seen in Table 3. Average value of absolute growth in length of each treatment was 1.59 cm; 1.61 cm; 1.58 cm; 1.21 cm; 1.55 cm; and 1.54 cm, respectively. Based on the analysis of One-Way ANOVA on absolute growth in length, a value of 0.744 (P>0.05) was obtained. Therefore, it is said that GFP gene did not significantly affect the absolute growth in length of betta fish.
As a supporting parameter in this study is the growth of larvae. Analysis result of growth rate in weight is presented in Table 3. Average growth rate in weight of each treatment was 5.39% day⁻¹; 5.53% day⁻¹; 6.04% day⁻¹; 5.86% day⁻¹; 6.33% day⁻¹; and 6.16% day⁻¹, respectively. Based on One-Way ANOVA, the growth rate in weight was 0.326 (P>0.05). Therefore, it is said that GFP gene did not affect the growth rate in weight of betta fish.

4. Conclusion

The method of transfection and electroporation was successfully used in GFP gene transfer in betta fish embryo. DNA concentration in GFP gene transfer by electroporation in betta fish embryo was 30 µg µL⁻¹.

References


reliably express GFP in whole muscles or the whole body by using promoters of zebrafish origin. *Dev Biol*. 192(2):289-299.


