Identification of sex-linked molecular markers in Indonesian giant freshwater prawn *Macrobrachium rosenbergii*

**ABSTRACT**

Male giant freshwater prawn grows faster than its female. Therefore, male mono sex culture is one of the solutions to improve aquaculture production. The all-male population of giant freshwater prawns can be produced by mating the neo-females (sex-reversed males) with the normal males. This study was aimed to identify the molecular markers related to the giant freshwater prawn sex. Specific primers were designed based on female-specific AFLP marker sequences to distinguish male and female sex on the prawns. Three locations for obtaining the Indonesian prawns in this study were Aceh, Sukabumi, and Solo. Based on the PCR analysis with MrMKn primers, 30 samples of female prawns had 100% occurred DNA bands, while no DNA bands were obtained in all-male prawns from Solo. Nevertheless, MrMKn primers still detected 10–16% male prawns from Sukabumi and Aceh. This indicated that MrMKn primers could not yet distinguish the male prawns for all populations. Moreover, the results suggested that the three prawn samples were different based on female-specific gene sequence. The MrMKn primers have the opportunity to be used in the selection of the female ZZ (neo-female) prawns from Solo without progeny test, so that the determination of female ZZ candidates can be identified more quickly. However, the primer still needs to be redesigned to distinguish neo-female prawns from Sukabumi and Aceh.

**Keyword:** giant freshwater prawn, mono sex, neo-female, sex markers

**ABSTRAK**

Udang galah jantan lebih cepat tumbuh dibandingkan dengan betinanya sehingga budidaya udang galah monoseks jantan menjadi salah satu solusi untuk meningkatkan produksi budidaya. Populasi monoseks jantan udang galah dapat dihasilkan dengan mengawinkan *neofemales* (*sex-reversed males*) dengan jantan normal. Sistem kromosom pada udang galah berbeda dengan ikan. Individu betina bersifat heterogametik (WZ) dan jantan homogametik (ZZ). Dalam perkembangannya, terdapat kendala dalam menentukan individu *neofemale* yang memiliki kromosom ZZ. Berdasarkan pendekatan sistem kromosom tersebut, maka dapat dijadikan acuan untuk membuat marka molekuler terkait kelamin udang galah. Penelitian ini bertujuan mengidentifikasi marka molekuler terkait jenis kelamin pada udang galah. Primer spesifik didesain berdasarkan sekuen *female specific AFLP marker* untuk membedakan kelamin jantan dan betina pada udang galah. Tiga sumber udang galah digunakan dalam penelitian ini, yaitu Aceh, Sukabumi, dan Solo. Berdasarkan hasil analisis PCR dengan primer MrKKn, dari 30 sampel pada kelompok udang galah betina diperoleh hasil 100% pita DNA muncul, dan tidak terdapat pita DNA pada semua udang galah jantan asal Solo. Namun demikian, primer MrMKn tersebut masih mendeteksi sebesar 10–16% pada udang galah asal Sukabumi dan Aceh. Hal ini menunjukkan bahwa primer MrMKn belum dapat membedakan udang galah jantan dari semua populasi. Selain itu, dapat dikatakan bahwa ketiga udang galah uji adalah berbeda, khususnya sekuen gen spesifik betina. Primer MrMKn berperlu digunakan dalam proses seleksi udang galah betina ZZ (*neofemale*) asal Solo tanpa harus melalui uji progeni sehingga penentuan kandidat betina ZZ lebih cepat teridentifikasi. Akan tetapi, primer masih perlu didesain ulang untuk membedakan *neofemale* asal Sukabumi dan Aceh.

Kata kunci: marka kelamin, monoseks, neo-female, udang galah
INTRODUCTION

Giant freshwater prawn *Macrobrachium rosenbergii* is one of crustacean species with high economic value. It has some excellences, its larger body size (Suwartiningsih & Utami, 2020) and its endurance against some of environmental factor, especially against salinity (Khasani, 2013). Due to the excellences of giant freshwater prawn caused vivid giant freshwater prawn culture in some countries in Southeast Asia, South Asia, and North Australia. Nowadays, giant freshwater prawn culture is massively developed in various countries, such as in United States of America and China (Akter et al., 2014; Zafar et al., 2015; David et al., 2018). In 2009-2012, the production of giant freshwater prawn has contributed over 57.1% of the total production of freshwater prawn *Macrobrachium* sp. globally (New & Nair, 2012). The annual production of giant freshwater prawn is known over 600,000 tons in 2020 (Maliwat et al., 2020). The northwestern of India, Vietman, Philippines, Papua New Guinea, and north of Australia are considered as main distributor in giant freshwater prawn production activities (Banu & Christianus, 2016; Maliwat et al., 2020).

The growth of male giant freshwater prawn is faster than the female, therefore the size of consumption is easy to reach (Sui et al., 2019; Tan et al., 2020). This attracts the aquaculturist to monosex culture of male giant freshwater prawn in order to get higher production of giant freshwater prawn with shorter rearing time (Nair & Salin, 2012). The mono sex breeding is considered more profitable, both in agriculture and the aquaculture sectors. The profits potential are offered various, i.e better growth performance and reduce the impact of territorial nature (Lezer et al., 2015). The monosex culture of male giant freshwater prawn produces better growth performance and better weight gain than the female one, thus are more profitable economically (Arisandi, 2013; Piyaviriyakul & Darawiroj, 2014).

The development of monosex culture of male giant freshwater prawn requires a sustainable supply of male giant freshwater prawn fry. Therefore, by mating the neo-females/ZZ (sex reversed-males) with the normal male, is one of the solutions to produce male mono sex of giant freshwater prawn (Aflalo et al., 2006; Aflalo et al., 2012). The sex chromosome of female giant freshwater prawn is heterogametic (Justo et al., 1991), with the chromosome code was WZ (Malecha et al., 1992). Furthermore, some of the previous studies has confirmed that there are other species with sex chromosome of WZ/ZZ, i.e., crayfish and white shrimp (Guo et al., 2019; Levy et al., 2020). Levy et al. (2020) stated that the chromosome system of WZ/ZZ is commonly found in macruran decapoda crustacea (crayfish and lobster). However, until now, there is no identification report about sex chromosome mating of giant freshwater prawn from karyotyping process.

Sex identification of female giant freshwater prawn/ZZ could be determined visually and confirmed by progeny testing, yet those steps need longer time and higher cost. Thus, the PCR method found potentially to accelerate the identification of neo-female giant freshwater prawn. The study about the relationship of sex and molecular marker is less reported. Moreover, the further analysis of nucleotide sequences has shown that this marker is specific female alleles (Jiang & Qiu, 2013). Currently, there has been no verify the sensitivity of the marker that reported by Jiang and Qiu (2013) in the giant freshwater prawn in Indonesia. According to those informations, this study is done to identify sex-linked molecular of giant freshwater prawn by using specific primer, allowing to identify the sex in early development stage of giant freshwater prawn when the phenotype differentiation is yet unidentified. The aim of this study was to verify the sensitivity of MrMK primer and to obtain specific primer that may to distinguish the sex of three populations of giant freshwater prawn in Indonesia. The result of this study is expected can be used to identify the sex in early development stage of giant freshwater prawn and is expected can be applied in the next individual selection stage as the result of sex reversal, thus to minimize selection time.

MATERIAL AND METHOD

Tested fish

Tested fish that used in this study was the population of male and female giant freshwater prawn obtained from Aceh (Sungai Peurlak) and Solo (Sungai Bengawan Solo), and male and female giant freshwater prawn of SIRATU variety. Giant freshwater prawn of SIRATU variety comes from three different varieties, i.e., from Sungai Bone, Sungai Mahakam, and Sungai Citanduy with nine crossbreed combinations. The individual selection was done in Pelabuhan Ratu therefore this giant freshwater prawn is called...
SIRATU giant freshwater prawn, as mentioned in A Decree of Ministry of Marine Affairs and Fisheries Republic of Indonesia number 25/ Kepmen-KP/2015 on April 16th, 2015. The giant freshwater prawn weighed about 50 to 60 grams/ individual.

All populations of giant freshwater prawn were directly acclimated at less than 60 mins. After that, the giant freshwater prawn was put into a rearing basket sizing of 30 cm × 20 cm × 10 cm with 1 individual/rearing basket in order to minimize the cannibalism since the giant freshwater prawn was still weak. The rearing baskets were put into a fiber tank sizing of 240 cm × 150 cm × 50 cm with a volume of 1.8 m³. The water level of each rearing tank was between 30–40 cm with aeration system, water inlet, and outlet pipe. Female and male giant freshwater prawn were reared in rearing tank separately. The water was siphoning off every day, to remove the sediment of waste in the bottom of the tank. The tank was cleaned up once in couple days by draining the water, brushing the tank, and cleaning the feces and feed waste, therefore the hygiene of the tank and the water quality remain maintained.

The feed was commercial feed types pellet food with high protein, squid, and tubers. The feeding frequency was four times a day. Squid and pellet feeding was given at daytime. Meanwhile, at the evening, the giant freshwater prawn fed with sweet potato. As many as 30 male and female giant freshwater prawns from three populations were collected to perform the DNA genome extraction. The giant freshwater prawn that used for this study were already experienced sex differentiation. The sex was differentiated based on Aflalo et al. (2006).

**Genome extraction**

The genome extraction method was based on Sambrook manual book (1989). The DNA extraction was performed using pleopods pieces. A phenol chloroform method was used for DNA extraction. The extraction sample can be stored in refrigerator with temperature of -20°C until it is ready to be used.

**The verification of MrMK primer sensitivity**

The verification of MrMK primer sensitivity that designed by Jiang and Qiu (2013) was used nine giant freshwater prawns from Sukabumi. The primer of MrMK sequences are forward 5’-GTGCGTGACATCAAGGAA-3’ (19 bp) and reverse 5’-GAATGGCTCTCGTGAATGC-3’ (19 bp). As an internal control, the DNA loading was β-actin primer (forward 5’-GATGAGTCCTGAGTAACAA-3’ and reverse 5’-TAAGTGGTTCTCGTGAAATGC-3’). Amount 1 µg of DNA was used as PCR sample, then it mixed with 1 µL of forward and reverse primer (10 pmol), 5 µL of MyTaq Redmix (Bioline, USA), and it added with nuclease-free water until the final volume reached 10 µL. The PCR process was run in 95°C for five mins, 1 cycle; (95°C for 30 sec; 49,5°C for 35 sec; 72°C for 30 sec), 35 cycles; 72°C for 2 mins, 1 cycle. Amount 3 µL of DNA product of PCR amplification was used for electrophoresis using 1% agarose gel and DNA marker. The visualization of DNA bands was performed by using GelDoc and ultra violet light.

**DNA purification**

The DNA fragment as the result of PCR amplification was purified from agarose gel to minimize the DNA contamination while DNA ligation. The PCR result was electrophoresis by using 1.5% of agarose gel, then the DNA was isolated from the gel by using a Geneaid™ Gel kit or PCR DNA Fragments Extraction kit (Geneaid, UK). The gel was cut on the part of DNA bands along with the target length, was 700 bp and was put into microtube. Amount 3 µL as the result of DNA that isolate from the gel was electrophoresis by using 1% of agarose gel to confirm the success of DNA extraction from the gel.

**The DNA ligation and transformation**

The DNA fragment as the result of gel purification was connected with pGEM-T Easy cloning vector. The compositions of ligation reactor were 5 µL of DNA solution, 0.5 µL of pGEM-T Easy, 6.5 µL of 5x ligation buffer, and 1 µL of T4 DNA ligase enzyme (TAKARA). The incubation was performed for two hours at room temperature, then was incubated overnight in the chiller at 4°C. Amount 6.5 µL of the result of ligation reaction was mixed into the microtube with competent cells. The DNA transformation was performed by using heat shock at 42°C for 60 sec. After that, the sample was incubated on-ice for two to three mins, then 900 µL of SOC solution was put into the microtube. Afterward, the microtube with bacteria of the transformation result was incubated at 37°C for an hour. Then the bacteria were spread on the petri disc SOB agar media contained IPTG, X-gal, and ampicillin, and was incubated at 37°C, overnight.

The white colony that grew on the SOB agar
media was taken up by using a steril toothpick and was put into a test tube with SOB media contained 100 μL/mL of ampicillin. The plasmid bacteria colony was isolated by using an alkaline lysis solution mini preparation technique (Sambrook & Russell, 2001).

The verification of the result of plasmid isolation
The verification of the result of plasmid isolation was performed by using PCR method to find the DNA insertion. The examination of DNA insertion was performed by using MrMK primer. The PCR process was run at 94°C for five mins in only one cycle; (94°C for 30 sec; 51.5°C for 30 sec; 72°C for 30 sec) in 35 cycles; 72°C for two mins in one cycle; and 4°C (unlimited). The result of positive insert PCR was clearly marked by the appear of 700 bp of DNA bands.

Sequencing and MrMK specific primer design
The plasmid that contained DNA fragment of PCR amplification product with MrMK primer, then was sent into Firstbase Sequencing, Singapore. The MrMK primer still not worked successfully to determine all the population of giant freshwater prawn, therefore the sequencing and new primer design are still needed. The sequencing result should be aligned to find out the different area to be used for specific primer. The sequences from male and female giant freshwater prawn were aligned with BIOEDIT program ver. 7.2. Furthermore, from the alignment result, the sequence was selected (only a pair of forward and reverse primer as specific primer) for all three populations of giant freshwater prawn. The PCR optimization was done by modifying the annealing temperature, allowing the specific PCR product for female giant freshwater prawn.

Data analysis
The analysis data of isolation result and characterization of sex marker in the giant freshwater prawn was performed descriptively. The sequencing data obtained from this study, then compared with the identical level of the genome of giant freshwater prawn which has been available in the Gene Bank by using basic local alignment search tool (BLAST) software in NCBI (http://blast.ncbi.nlm.nih.gov/Blast.cgi). The trace and the alignment of sequencing result and the making of new primer design used BioEdit program, MEGA 6 (Tamura et al., 2013). The primer that was specific and accurate could determine the giant freshwater prawn in all populations.

RESULT AND DISCUSSION

Result
The verification of MrMK primer sensitivity
First, the verification of MrMK sensitivity primer was performed. The PCR result used forward dan reverse of MrMK primer showed that the product is clearly seen with DNA length bands was 700 bp and the β-actin was around 250 bp (Figure 1). Afterward, that primer was tested with all the samples of giant freshwater prawn from Aceh, Sukabumi, and Solo. The recapitulation of PCR testing result showed in the Table 1 below. All the samples of female giant freshwater prawn from three different locations have been detected with DNA length bands was 700 bp, meanwhile there still was 16% of male giant freshwater prawn from Solo and 30% of each giant freshwater prawn from Aceh and Sukabumi. By this, the further step, new primer is designed.

![Figure 4. Nitrogen output distribution in biofloc-based C. gariepinus culture at different densities (500, 750, and 1000 fish/m³) and a control after 8 weeks of culture.](image-url)
Sequencing, new primer design, and sensitivity test

The PCR product of female giant freshwater prawn measuring 700 bp of DNA bands length was cloned to cloning vector. Then the plasmid used for sequencing. This study used three different colonies for the sequencing. The sequencing result of PCR product and its alignment showed in Figure 2. The sequencing result showed similar in all plasmid samples from bacteria colonies. From this alignment result, the sequence of male and female giant freshwater prawn obtained different area used to design a specific primer (it marked with a box and an arrow).

One set of specific primer candidate (MrMKn) used as a molecular marker as sex distinction in designated male and female giant freshwater prawn (Figure 2), i.e., forward MrMKn5’-CAGTATTTCCGGAATGATTTGCTCGGG-3’ and reverse MrMKn 5’-CCGATAACTCTCGCG-

AATGAGC-3’ with 390 bp of DNA bands target. The MrMKn primer was tested on three different populations of male and female giant freshwater prawn. The result of this study showed that MrMKn primer was completely specific in female giant freshwater prawn with DNA band showed 390 bp (Figure 3).

According to PCR analysis results, all the population of female giant freshwater prawn from Aceh, Sukabumi (SIRATU), and Solo showed the DNA bands (Table 2). The DNA band expected did not appear in all populations of male giant freshwater prawn, yet this DNA band was unfortunately appeared in the population of male giant freshwater prawn from Aceh and Sukabumi, which each of them were 5 (16%) and 3 (10%). This showed that the sensitivity of MrMKn to determine the male and female giant freshwater prawn from Solo, SIRATU, and Aceh were 100%, 90%, and 84%.

| Male | Female |
|------|--------|
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |
| TTTGCCGATCGCTGGTATGCGGTATCGCTGGCGTTGCGT |

Figure 2. The alignment of DNA sequences of male and female giant freshwater prawn as the result of sequencing with MrMK primer.

| Male | Female |
|------|--------|
| 6917 |
| 763 |
| 977 |
| 803 |
| 1037 |
| 863 |
| 1097 |
| 923 |
| 1157 |
| 983 |
| 1217 |
| 1043 |
| 1277 |
| 1103 |

Figure 3. The PCR amplification with MrMKn primer in female giant freshwater prawn with DNA bands showed 390 bp (up), and the β-actin was 250 bp (bottom). The arrow at the right side showed the PCR product target. M= DNA marker; 1–12= the number of genome sample of giant freshwater prawn.
Discussion

Jiang and Qiu (2013) have already succeeded in obtaining specific primer of sex markers in the giant freshwater prawn in China. Furthermore, the MrMK primer has already verified its sensitivity in three different populations of giant freshwater prawn in Indonesia, yet the MrMK primer still could not determine specifically between male and female individuals, and not all the samples of giant freshwater prawn had a PCR product (Table 1). Therefore, the PCR product from male and female giant freshwater prawn were purified, then were sequenced to obtain the area that has selected as new specific primer target. The PCR product purification was intended to eliminate impurities in samples because it can disrupt the processes, i.e., DNA ligation, cloning, labelling sequencing, PCR, and RTPCR (Santo et al., 2017; Dwiyitno et al., 2018).

Table 1. The sex identification of giant freshwater prawn used MrMK primer in male and female giant freshwater prawn from Aceh, Sukabumi (SIRATU), and Solo.

| Sample | Sex | Positive PCR product (individual) | Percentage (%) |
|--------|-----|-----------------------------------|----------------|
| Aceh   | Female | 30                               | 100            |
| SIRATU | Female | 30                               | 100            |
| Solo   | Female | 30                               | 100            |
| Aceh   | Male | 9                                | 30             |
| SIRATU | Male | 3                                | 10             |
| Solo   | Male | 5                                | 16             |

According to the alignment data of sequencing result, one set primer was selected to be a sex marker of giant freshwater prawn from Indonesia. That area was selected because the 3 end was played important role when annealing and primer extension when PCR started (Yustinadewi et al., 2018). Afterward, the nucleotide bases G and C that had three hydrogen bonds was remained stable than nucleotide bases A and T that only had two hydrogen bonds (Poater et al., 2014; Nur’aini et al., 2019). The PCR process involved various steps, i.e., denaturation, primer annealing, and primer extension (Porta & Enners, 2012; Fitriatin & Manan, 2015). The optimal annealing temperature annealing should be noticed. The temperature optimization for annealing process was made for the primer to attach better, to multiply the PCR product, and to increase DNA specificity (Kurniawati & Hartati, 2018). The MrMKn primer in this study has DNA bands length of 390 bp. The optimization result showed that the best annealing temperature was 43°C, meanwhile Walker and Rapley (2002) stated that the recommendation of annealing temperature is between 40–60°C. The error of primer attachment can occur when the annealing temperature is too low. Yet, some of primers also can not attach well when the temperature is significantly increased (Naqib et al., 2019).

Based on PCR analysis, within 30 samples of giant freshwater prawn from Solo, the DNA bands were showed 100%, and the DNA bands were not occurring in all male giant freshwater prawn (Table 2). However, the MrMKn primer sensitivity in the giant freshwater prawn from Aceh was 84% and from SIRATU was 90% (Table 2). It showed that MrMKn primer still could not determine male giant freshwater prawn in all populations. In the early stage, the primer was designed by the result from BLAST and was specific. Nevertheless, the lack of the information about the genome sequence of giant freshwater prawn in Indonesia could complicate to collect specific primer of all populations. Moreover, it concluded that the giant freshwater prawn used in this study was totally different, especially the genome sequence of sex. The MrMKn primer can be used in the selection process of ZZ female giant freshwater prawn from Solo without progeny test, therefore the determination of ZZ female candidate can be faster. Yet, the primer is still needs to redesign to determine the neofemale from Sukabumi and Aceh. The further study can be performed by isolating the whole sequences to obtain other candidate from different areas to determine some population of giant freshwater.
prawn in Indonesia. Some studies showed that there is a potency that several sex genes can be used as markers. Yu et al. (2014) explained that the result of qPCR analysis of Dmrt genes in the giant freshwater prawn showed a different gene’s expression pattern in the development of sexual dimorphism and the embryo, i.e., MroDmrt11E and MroDmrt99b gene. Furthermore, the content of N-acetyltransferase and melatonin in optic lobe of male giant freshwater prawn was not significantly different, yet in female giant freshwater prawn was otherwise (Withyachumnarnkul et al., 1999).

CONCLUSION

This study proved that the MrMKn primer could determine ZZ male and ZZ female of giant freshwater prawn from Solo specifics, yet it could not determine the giant freshwater prawn from Aceh and Sukabumi (SIRATU). The further study should be performed to obtain specific primer that can determine all populations of giant freshwater prawn in Indonesia.

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