The Development and Application of Genetic Markers for Giant Freshwater Prawns *Macrobrachium Rosenbergii* by Microsatellites

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**Abstract**

This article reports the development of microsatellite primers for giant freshwater prawns (*Macrobrachium rosenbergii*), using the genome library for this type of prawns and biotinylated probe in isolating six types of DNA fragments with base sequence repeats, namely (AG)10, (TG)10, (CAA)10, (CAG)10, (GAT)10 and (TAC)10. Four clones harboring microsatellites were chosen, namely SH2-9F, SH2-10C, SH2-11D and SH3-11G. Base sequencing found no microsatellite sequences. Four pairs of primers were designed, namely DTLSSH 7, DTLSSH 8, DTLSSH 9 and DTLSSH 12, respectively. These primers were tested on the DNA of giant freshwater prawns, and DNA band sizes were found to be 131, 174, 210 and 193 per 205 bp, respectively. The developed microsatellite primers may be used in conjunction with primers for other types of prawns. Genetic diversity checking of giant freshwater prawns in Thailand showed that their population in Thailand may be classified into two groups: one with the closest genetic relationship being those from Khon Kaen, Samut Songkhram and Ang Thong provinces while the other with similar genetic characteristics were those from these three provinces as compared to those from Surat Thani province.

**Keywords:** Giant freshwater prawn; Microsatellite; Genetic marker

**Introduction**

*Macrobrachium rosenbergii*, or its generic name the giant freshwater prawn, is the freshwater prawn popularly raised in India, in the Southeast Asian countries, in Northern Australia and in certain parts of the Pacific and Indian Oceans. In Thailand, the aquaculture part is considered an industry with great significance to the country as prawns can be a principal aquariumal protein source that is high in nutritional values. In addition, they are one of Thailand’s top export goods in terms of income earnings. Their domestic productivity enhancement, however, is without stability, and there are some limitations in their production. Augmented growth rates of prawns would be another indicator of productivity potentials. This condition partly requires improvement in genetic characteristics [1]. Selection of suitable stocks in breeding programs is affected by genetic variation of any given animal population. Development of microsatellite markers will, therefore, be very useful in identifying polymorphisms and even in the mapping of a sample population from the same species. The polymorphisms of these markers come from their different numbers of repeats. Microsatellite markers are codominant-hence can detect the difference between homozygotes and heterozygotes. At present, microsatellites in giant freshwater prawns have been developed from two groups; one of which is the eastern form [2] with six pairs of primers having been developed [3]. The other group is the western form [2]. Thailand’s giant freshwater prawns have been classified in this latter group, which has so far had 11 pairs of microsatellite primers [4]. These, however, are not sufficient for use as molecular markers for giant freshwater prawns in Thailand. Its data base for development of additional microsatellite primers to cover other parts of the prawns’ genome would maximize the usage and accuracy in the breeding programs in the future.

**Material and Methods**

1. Collection of sample stocks of giant freshwater prawns general in Thailand, DNA extraction of the prawns (*Macrobrachium rosenbergii*) and DNA quantification, using electrophoresis.
2. Digestion and ligation of liker to DNA fragments. The extracted DNA or genome was digested, using a restriction enzyme as follows: Approximately 500 ng of the genomic DNA were digested using the restriction enzyme Tru 9I or MseI and ligated using T4 ligase and MseI-Adaptor at 37°C for seven to eight hours or overnight. After the incubation and digestion with the restriction enzyme, the DNA was then made 10-20-fold diluted in Ultrapure water.
3. Capture of specific repetitive sequences with biotinylated oligonucleotides. Probes were constructed from base sequences with multiple repetitions at the same locations, such as (AG)10 or (C-AA)10, as follows: Utilized probes consisted of six types of biotinylated oligo SSR, namely (AG)10, (TG)10, (CAA)10, (CAG)10, (GAT)10 and (TAC)10. The oligo SSR contained conjugated biotin at the 5’ terminal of each strand. The probe/DNA fragments ratio during probe preparation prior to hybridization was 10 pmol/1.0 pmol. Subsequently, each biotinylated oligo SSR was mixed with 1 ml of Streptavidin Magnetic Sphere Paramagnetic Particles (SA-PMFs). The streptavidin binds to biotin while its other end binds to magnetic particles. The mixture was left at room temperature for two hours and then washed with 5X SSC twice while using Magnetic Sphere Magnetic Separation Stands (MS-MSS) to hold the biotinylated oligo SSR bound to the streptavidin. Excessive biotinylated oligo SSR was removed by washing, and the desired part was subsequently hybridized with the DNA.

a. In mixing the DNA with the streptavidin-biotinylated oligo SSR complex, the former was first separated into single strands, applying heat at 95°C for 10 minutes. It was immediately put on ice and then mixed with SSC at a concentration adjusted to 6X. The temperature

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used was dependent on the probes’ melting temperatures: 55°C for (AG)10 and (TG)10, and 65°C for (CAA)10, (CAG)10, (GAT)10 and (TAC)10. The reaction was allowed to take place for approximately 10-12 hours or overnight. Subsequently, only the DNA that had been bound to streptavidin-biotinylated oligo SSR complex was selected, using MS-MSS to draw it to the sides of tubes, and the supernatant removed. The next step was to wash with low and high stringency solutions to remove any remaining unwanted DNA and to clean the desired DNA. A subsequent step was to separate the latter from the streptavidin-biotinylated oligo SSR complex by adding 0.15 M NaOH and leaving the mixture at room temperature for approximately 30 minutes. MS-MSS was then used to draw the streptavidin-biotinylated oligo SSR complex to the sides of tubes. All supernatant was transferred to new tubes and neutralized by adding 1 N acetic acid and TE buffer. The obtained DNA was subsequently washed clean, using High Pure PCR isolation kit (Boehringer).

4. Library construction and cloning, using pGEM-T Easy kit (Promega) as follows:

a. Vector used in the ligation to the DNA was pGEM-T Easy (Promega) sized approximately to 3 kb at the concentration of 50 ng/µl. The ligation step utilized the insert/vector ratio of 3:1 with the following formula in calculating the quantity of insert to use.

\[
\frac{\text{Insert size (kb) x vector quantity}}{\text{Vector size (kb)}} = \text{Insert quantity to use}
\]

b. A total 10 µl of ligation reaction comprised 25 ng of pGEM-T Easy, X ng of insert, 5 µl of 2X Rapid Ligation buffer and 3 µss units of T4 ligase. The reaction was left at 4°C overnight. The next step of transformation employed electroporation, using E. coli strain DH 10B as host and plating on selective agar medium (100 µg/ml ampicillin, 100 µM IPTG and 40 µg/ml x-gal). The obtained white colonies were checked for the insert size, using PCR with primers M13 forward and M13 reverse. Electrophoresis was used to compare with control, which consisted of a blue colony, and molecular weight markers. Only white colonies with inserts sized 500 bp and larger were chosen for the sequencing step.

5. Screening of positive colonies by PCR

a. Dot blot hybridization of oligonucleotide probes as per the method described by Sambrook [5].

6. DNA sequencing of positive clones:

7. The desired colonies were chosen to be cultured in a broth containing 1 mL of ampicillin (100 µg/ml) at 37°C for approximately three to five hours. The cell culture underwent PCR, of which 20 µl comprised 0.25 pmol of M13 primers (forward and reverse), 100 µM dNTPs, PCR buffer (100 mM Tris-HCl (pH 9.0), 500 mM KCl, 1.5 mM MgCl2) and 1 µTaq DNA polymerase, in GeneAmp PCR System 9700 (Applied Biosystems). Its programmed temperatures were as follows: one round of 94°C for three minutes, 94°C for 30 seconds, 55°C for 30 seconds, 35 rounds of 72°C for two minutes each round and one round of 72°C for seven minutes. After the reaction was complete, the PCR product concentrations and DNA base sequences were determined.

8. Sequencing analysis, using dye terminator cycle sequencing and ABI PRISM 377 DNA sequencer (PE Biosystems).

9. Primer design was based on the determined base sequences, using Gene Fisher interactive primer design software or Primer 3 in designing primers for use in DNA amplification. The SSR part was designed based on flanking region of that SSR strand, taking into account the length of oligonucleotides, the melting temperature and % G+C content. In addition, primers can be designed using such computer software as Primer Express (Perkin-Elmer), Prophet and also online design software. For example, website of http://www.genome. wi.mit.edu/cgi-bin/primer/primer3.cgi/ can check characteristics of primers that have been designed by calculation on http://www.genosys.com/. In the primer design, however, it is required that the location to be designed by the software should be quite far away from the SSR part; that is, approximately 30-50 bases away. Some primer may be less far away, but the distance should not be shorter than the primer length, and the base repeats found at that location should not be of more than three bases, particularly A and T. Primers were synthesized after obtaining the desired DNA base sequences for use the in primer design for the subsequent amplification of DNA, with base repeats as shown in Figure 1. The desired sequences comprised two short DNA strands; each of which was 18-25-base long and contained approximately 35-60% GC. In addition, each strand had to have a melting temperature between 55-65°C.

10. Primer testing and optimization.

Results

Genomic DNA preparation

The results of DNA extraction from giant freshwater prawns are demonstrated in Figure 2. Comparing with lambda DNA, which was used as a marker, the DNA concentration (SH) was approximately 50 ng/µl, which were sufficient for test in the next step.

DNA (100, 300, 500 and 1,000 ng, respectively); SH, DNA extracted from prawn samples

DNA fragments preparation and adaptor-ligated DNA

After the genomic DNA of the prawns was digested with the restriction enzyme Tru 9I or MseI and ligated with MseI-Adaptor, it was found that the prawn DNA samples were on all three A-lanes. The enzyme was able to digest the DNA completely, as illustrated in Figure 3.

Subsequently, DNA was amplified, using PCR technique by first making the ligated DNA approximately 10-20-fold diluted to 10-20 µl. In the PCR, the initial DNA was able to bind to the adaptor. One base

\[
\chi = \frac{X}{100}\%
\]

Figure 1: Locations on DNA strands for use in primer design.

Figure 2: Quantification of DNA extracted from giant freshwater prawns: M1-M4, lambda.
for use in selection was added to the 3’-end. Figure 4 shows the results of DNA amplification as seen by the more intense color bands on lane A than those in Figure 3.

**Enrichment**

Using six types of biotinylated oligo SSR probes, being (AG)10, (TG)10, (CAA)10, (CAG)10, (GAT)10 and (TAG)10. This step enabled DNA enrichment.

**Amplification**

The DNA obtained from the enrichment step was single-strand DNA, which could not be ligated into plasmid. The DNA, therefore, had to go through PCR, using primer MseI-5’GAT GAG TCC TGA GTA ANNN3’, to become double-strand DNA. The amplified DNA was to be precipitated, using 1/10 V NaOAc (pH 5.2) and 2 V 95% EtOH, left at -80°C for approximately one hour and centrifuged at 14,000 rpm for 20 minutes. The precipitate was washed with 70% EtOH twice, air-dried at room temperature and reconstituted with water. The resulting solution was measured in a spectrophotometer for use as insert DNA in the subsequent step.

**Library construction**

The prawn genome library was constructed for use in screening. The obtained white colonies were checked for the insert size. Electrophoresis was utilized to compare with the control, which was a blue colony, and molecular weight markers. Only the white colonies with inserts sized 500 bp and larger were chosen for the sequencing step.

**Sequencing and primer design**

At this stage, four clones were found that were expected to harbor microsatellites and thus checked and sequenced. All four clones contained repetitive base sequences that did not match the desired microsatellites, as demonstrated in Figure 5. Primers were designed, based on the location with the most repeats. Starting from clone 1, these were clone 1: SH1-9F, primer designation: DTLSH 7; clone 2: SH2-10C, primer: DTLSH 8; clone 3: SH2-11D, primer: DTLSH 9 and clone 4: SH3-11G, primer: DTLSH 12. The base sequences of all four pairs of primers are described in Table 1. The developed primers are, nevertheless, considered an initially essential and important tool to be investigated into their application in further biotechnological basic research as regards prawns. The obtained primers are considered very appropriate for use in the research on prawns as they have been developed from the genomic part of giant freshwater prawns.

**Primer testing**

The synthesized primers were tested with the following DNA: first, DNA of giant freshwater prawns (genomic DNA) used in the initial step, as a positive control. Secondary, DNA of unknown giant freshwater prawns and finally, DNA of giant tiger prawns.

Ten µl of reaction comprised 1 and 5 ng of starting DNA, forward and reverse primers of 0.5 µM each, 1X PCR buffer (100 mM Tris-HCl (pH 9.0), 500 mM KCl), 2 mM MgCl2, 200 µM dNTPs and 0.2 unit of Taq DNA polymerase. It was put in a pre-set temperature controller (GeneAmp® PCR System 9700; Applied Biosystems), using the following temperatures: one round of 94°C for three minutes, 94°C for 30 seconds, 60°C for 30 seconds and 40 rounds of 72°C for one minute per each round and one round at 72°C for five minutes, as detailed in Tables 2 and 3. The PCR results were checked, using 4.5% polyacrylamide gel followed by silver staining, and primer test results were obtained in Figure 6.

**Discussion**

Application in the Classification of Giant Freshwater Prawn Population in Thailand, giant freshwater prawn samples were collected from four provinces in Thailand, namely Khon Kaen (KK), Samut Songkhram (SK), Surat Thani (SR) and Ang Thong (AT). Twenty-five prawns were collected from each reservoir, totaling 100 samples. Data from the study on genetic diversity of giant freshwater prawns were analyzed by applying genetic distance values to the creation of a dendrogram among prawn population groups in Thailand by using UPGMA (unweighted pair-group method arithmetic means) [6] and NTSYSpc Version 2.10p. The giant freshwater prawn population in Thailand could be classified into two groups (Figure 7): one with the closest genetic relationship, namely those from Khon Kaen, Samut Songkhram and Ang Thong provinces, and the other with similar genetic characteristics, namely those from these three provinces as compared to those from Surat Thani province. This classification corresponds to Daungwonsga [7], who reported that the giant freshwater prawn population group from Khon Kaen was genetically close to that from Suphan Buri, a province in the Central region like Samut Songkhram and Ang Thong, and that the groups from Khon Kaen and Suphan Buri were genetically different from that from Surat Thani. The findings of the present study are consistent with the geographical distance; that
is, Samut Songkhram is closest to Ang Thong, and both of them are in the Central region. The reason that the prawn population from Khon Kaen has similar genetic characteristics to that from Samut Songkhram and Ang Thong may be that the prawn breeding or offspring has been moved from the Central region areas for the aquaculture in Khon Kaen area and has then escaped from culture houses or farms to natural reservoirs as well as the fact that the prawns have been allowed to live in natural reservoirs in order to increase productivity. Likewise, the prawn population from Khon Kaen has more genetic similarity to that from Surat Thani than that from Samut Songkhram and Ang Thong. This similarity is best understood by considering the genetic distance values as shown in Table 4, which would not seem to correspond to the longer geographical distance between Khon Kaen and Surat Thani as compared to the distance between Samut Songkhram and Ang Thong.

This classification differs from what results from the use of apparent band data of microsatellite DNA in the classification since DNA is a component present in almost all cells in the same quantity. DNA from any tissues, growth phases or physiological conditions can, therefore, be examined without being dependent on the environment [9]. This approach provides greater accuracy than the population classification by morphology.

<table>
<thead>
<tr>
<th>No.</th>
<th>Primer name</th>
<th>Clone</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>DTLSH 7</td>
<td>SH2-9F</td>
<td>AAGCAAAACCAAAATATCTTACC</td>
</tr>
<tr>
<td>2</td>
<td>DTLSH 8</td>
<td>SH2-10C</td>
<td>GTACGAGGATCTAAAGGTGG</td>
</tr>
<tr>
<td>3</td>
<td>DTLSH 9</td>
<td>SH2-11D</td>
<td>AATTGTTGCTCTGTCAGGG</td>
</tr>
<tr>
<td>4</td>
<td>DTLSH 12</td>
<td>SH3-11G</td>
<td>TCGCTGGAACACGGAATGG</td>
</tr>
</tbody>
</table>

Table 2: PCR reduction component and primer qualities.

<table>
<thead>
<tr>
<th>No.</th>
<th>Primer name</th>
<th>Annealing Mg Conc.</th>
<th>dNTPs</th>
<th>Cycle Number</th>
<th>Product Size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>DTLSH 7</td>
<td>60°C 2 mM 200 µM 40 cycles</td>
<td></td>
<td>131 bp</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>DTLSH 8</td>
<td>60°C 2 mM 200 µM 40 cycles</td>
<td></td>
<td>174 bp</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>DTLSH 9</td>
<td>60°C 2 mM 200 µM 40 cycles</td>
<td></td>
<td>210 bp</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>DTLSH 12</td>
<td>60°C 2 mM 200 µM 40 cycles</td>
<td></td>
<td>193/205 bp</td>
<td></td>
</tr>
</tbody>
</table>

Figure 5: DNA sequencing in clone 1-4 of primer design.

Figure 6: Bands of DNA from PCR products obtained from giant freshwater prawn samples by using the SSLP marker.
In the present development of microsatellite primers in giant freshwater prawns for use in examining the genetic diversity of the prawns, four clones harboring microsatellites were chosen and used in designing four pairs of primers: DTLSH 7, DTLSH 8, DTLSH 9 and DTLSH 12. The primers was utilized to test positive control, unknown prawn samples 1 and 2 and also giant tiger prawn samples. All four primers were able to produce distinct DNA bands of PCR products and confirmatory bands of PCR products. Any pair of primers chosen can give the same readings. Apart from these four pairs, anyone wishing to study and examine the genetic diversity of giant freshwater prawns needs also to use primers developed from additional work by other researchers. Among these other researchers [4], who reported the development of 10 pairs of microsatellite markers in giant freshwater prawns in Thailand, thus obtaining results that are correct and matching the examining requirements. In addition, prawn samples to use in studies should be taken from a variety of sources.

The classification of giant freshwater prawn population in Thailand shows low-level genetic diversity with an average genetic distance of 0.165. The information on apparent bands of microsatellite DNA has classified the population of these prawns into two groups: one with the closest genetic relationship, namely those from Khon Kaen, Samut Songkram and Ang Thong, and the other with similar genetic characteristics, namely those from Surat Thani. These findings are consistent with the geographical distances.

**Conclusion and Suggestions**

In the present development of microsatellite primers in giant freshwater prawns for use in examining the genetic diversity of the prawns, four clones harboring microsatellites were chosen and used in designing four pairs of primers: DTLSH 7, DTLSH 8, DTLSH 9 and DTLSH 12. The primers was utilized to test positive control, unknown prawn samples 1 and 2 and also giant tiger prawn samples. All four primers were able to produce distinct DNA bands of PCR products at the base sequences of 131 bp (DTLSH 7), 174 bp (DTLSH 8), 210 bp (DTLSH 9) and 205 bp (DTLSH 12). Although the test with these four pairs of primers in different types of prawns yielded equal DNA band values, it is not clear whether they really are identical or different. Choosing and developing primers from each type of prawns are needed for a parallel test for comparison and confirmation. In designing these four pairs of primers in giant freshwater prawns, optimal conditions for their use were determined as demonstrated in Tables 2 and 3. The test results have indicated that the primers are specific to individual locations and yield distinct DNA bands of PCR products. Any pair of primers chosen can give the same readings. Apart from these four pairs, anyone wishing to study and examine the genetic diversity of giant freshwater prawns needs also to use primers developed from additional work by other researchers. Among these other researchers [4], who reported the development of 10 pairs of microsatellite markers in giant freshwater prawns in Thailand, thus obtaining results that are correct and matching the examining requirements. In addition, prawn samples to use in studies should be taken from a variety of sources.

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