Detection of Microcystin (mcyE) Gene in Recreational Lakes in Miri, Sarawak, Malaysia

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http://dx.doi.org/10.12944/CWE.11.3.02

(Received: November 22, 2016; Accepted: December 14, 2016)

ABSTRACT

Toxic cyanobacteria blooms became a worldwide problems as many countries encounter the presence of the blooms in most of water bodies. As part to develop monitoring of cyanobacterial toxins in Malaysia, samples taken in twelve points in five different lakes in Miri, Sarawak. Polymerase chain reaction (PCR) amplification of cyanobacterial 16S rRNA were carried out to detect the presence of cyanobacteria in the water samples. Cyanobacterial 16S rRNA were detected in all the samples collected. While molecular analysis for detection of cyanobacterial toxin encoding gene were done using specific primers. PCR amplification of cyanobacterial toxin-encoding gene were carried using the combination of forward primer; mcyE-F2 and reverse primer; mcyE-R4 to amplify generic microcystin (mcyE) gene in the samples. Out of twelve samples collected, microcystin (mcyE) producing gene was detected in one of the samples tested. Presence of microcystin encoding gene indicates the risk of cyanobacterial toxins in Miri, Sarawak.

Keywords: Toxic cyanobacteria, Cyanobacterial toxins, Cyanobacterial 16S rRNA, PCR amplification, microcystin.

INTRODUCTION

Cyanobacteria, or also known as blue-green algae are prokaryotes with unique characteristics. Unlike most prokaryotes, cyanobacteria consist of chlorophyll that enables them to obtain their nutrient mainly through photosynthetic action. Photosynthesis of the bacteria is important to provide atmospheric oxygen for other organism usage. Cyanobacteria are also important as a potential resource for renewable energy and natural products. However, excessive growth of cyanobacteria leads to formation of cyanobacteria blooms in the water environment. The blooms cause several problems such as unpleasant odour and taste, and most importantly, toxin production.

Capability of producing toxins by cyanobacteria was recorded earliest by Francis. However, understanding of the toxins structure, its genome code and biochemical pathway were known only after few decades later. Cyanobacterial toxins are encoded in a unique gene cluster, which is not species specific, but rather gene specific. That is, same species of cyanobacteria may or may not encode for the same toxin gene and that the same toxin may be encoded by several different species. The toxins are generally categorized in four major groups based on its toxicology effects, which are hepatotoxic toxin (microcystin and nodularin), neurotoxin (anatoxin-a, anatoxin-a(s), and saxitoxin), cytotoxic toxin (cylindrospermopsin) and dermatotoxin (aplysia toxins, lyngbyatoxin-A). Previous record showed that all the toxins are harmful to human and animals. In 1996, direct exposure to the cyanobacterial toxin during
routine hemodialysis treatment had caused 116 patients from Brazil to develop hepatotoxicity and neurotoxicity symptoms and 52 of them died due to the cyanobacteria toxification. The symptom is now can be referred as Caruaru syndrome⁶. Efforts in analysing health risk posed by cyanotoxin had been made several times and in 1999, World Health Organization (WHO) had included cyanobacterial toxins in the drinking water guidelines and recreational water guideline.

Currently, more than 65 countries worldwide including Thailand, Vietnam, Philippine, and Singapore have recorded the detection of toxic cyanobacteria in the water environment, with Malaysia confirming presence of toxic cyanobacteria in 2015⁷ and toxin-producing Microcystis was successfully isolated from Ayer Itam reservoir, Penang by Sim Yi Jing⁸.

As such, detecting toxin encoding gene cyanobacteria are crucial to clarify the status of cyanobacteria toxin in Malaysia. Researchers have established many toxic cyanobacteria detection methods. Most commonly used in detecting toxin encoding gene cyanobacteria involved molecular typing method such as conventional polymerase chain reaction (PCR). Using the PCR, presence of cyanobacterial toxin encoding gene in Miri, Sarawak, was investigated. To our best knowledge, this is the first research in detecting toxin gene encoding microcystin was conducted in Borneo Malaysia.

**METHODS**

**Sampling**

The locations involved a district in Sarawak, called Miri. Samples were collected in twelve points around Miri on June 2015. The lakes were located at Taman Tunku, Taman Awam Miri, Miri City Fan, Taman Bulatan and Taman Hilltop (Figure 1). In general, these lakes were manmade lakes located in residential and recreational areas in Miri, Sarawak. These sampling sites were chosen according to the visibility of the greenish organisms in the water bodies indicate the presence of the cyanobacteria. Accessibility and wind direction also taken into consideration in choosing the samples collection stations.

**DNA extraction**

DNA extractions from the environment were performed based on Bacterial DNA Extraction Kits (Vivantis) protocol. In brief, 1ml of cyanobacterial samples were centrifuged at 6000 rpm for 2 minutes and the supernatant were decanted completely. Then, 100µl Buffer R1 were added to the pellet and the cells were resuspended completely. 10µl of lysozyme were added into the cell suspension. The cell suspension were mixed thoroughly before incubated at 37⁰C for 20 minutes. After that, the digested cells were pelleted by centrifugation at 10,000 rpm for 3 minutes and supernatant were decanted completely. The pellet were resuspended in 180µl of Buffer R2 and proteinase K were added. The cells suspension were mixed thoroughly and incubate for 20 min in a shaking thermomixer. Two volumes of Buffer BG (approximately 400µl) were added and mixed by several times until a homogenous solution is obtained. The mixture were incubated for 10 min at 65⁰C before 200µl of absolute ethanol were added. The mixtures were then transferred into a column microfuge. The column was centrifuged at 10,000 rpm for 1 min. The flow through was discarded and washed using Wash Buffer⁹. The column was centrifuged for one minute at 10,000 rpm. The column was further centrifuged at the 10,000 rpm for 1 minute to remove residual ethanol. Finally, the extracted DNA was eluted into using pre-heated elution buffer.

Efficiency of DNA extraction was visualized by running through agarose gel. Extracted DNA was stored at -20 degree until required.

**PCR**

Cyanobacterial 16S rRNA were detected using primer pairs CYA106F, and CYA781R (a), CYA781R (b) and the sequences for the primer were shown in Table 1¹⁰,¹¹,¹².

PCR amplifications were performed with a Mastercycler® ep (Eppendorf). PCR reaction were carried out in 25 µL reaction mixtures containing 12.5 µL of 2X Taq Master Mix (Vivantis Technologies), 0.25 µL of each primers (Integrated DNA Technologies) and 2µl of DNA sample combined with sterile distilled H₂O to a volume of 25 µL of total reaction. PCR protocols to amplify cyanobacterial 16S rRNA
involved an initial denaturation for 2 min at 95°C; followed by 30 cycles, each consisting of 60 s at 94°C, 60 s at 60°C, and 60 s at 72°C; and a final extension of 7 min at 72°C. After the amplification, the products was loaded on 1% agarose gel that was prepared by adding 0.25 g of agarose (Vivantis Technologies) to 25 mL of 1 x TAE buffer. 2.5 µL of gel stain (TransGen Biotech) added into the hot agar. 2 µL of 6X loading dye (Vivantis Technologies) added into the PCR products. Gels were ran at 70 V for 40 minutes and viewed using Gel Doc™ XR+ (Bio-Rad).

Generic microcystin (mcyE) gene amplified using forward primer; mcyE-F2 combined with reverse primer; mcyE-R4. While the detection microcystin (mcyE) gene specific for *Microcystis* sp. used primer pair mcyE-F2 and mcyE-R8, microcystin (mcyE) gene specific for *Anabaena* sp. used primer pair mcyE-F2 and mcyE-12R and gene specific for *Planktothrix* sp. used primer pair mcyE-F2 and mcyE-plaR3. The sequences for the primers used in amplification for microcystin gene were shown in Table 2.

For mcyE gene-generic primers, PCR protocols was performed as followed: The first step was an initial denaturation step of 2 min at 95°C followed by 35 cycles of 30 s at 94°C, 30 s at 56°C and 60 s at 72 °C and a final extension of 10 min at 72 °C. PCR protocols for all three mcyE gene-specific primers involved of an initial denaturation for 3 min at 95°C. For microcystin (mcyE) gene specific for *Microcystis* sp., the initial denaturation step was followed by 25 cycles of 30 s at 94°C, 30 s at 60°C, and 60 s at 72 °C. For microcystin (mcyE) gene specific for *Anabaena* sp., the initial denaturation step was followed by 30 cycles of 30 s at 94°C, 30 s at 58°C, and 60 s at 72 °C. For microcystin (mcyE) gene specific for *Planktothrix* sp., the initial denaturation step was followed by 30 cycles of 30 s at 94°C, 30 s at 57°C, and 60 s at 72 °C. All the PCR cycles were followed by a final extension of 10 min at 72 °C.

**DNA purification**

DNA product from the PCR amplification were performed based on Ambiclean Kits (PCR & Gel) protocol (Vivantis Technologies) and sent to Center for Chemical Biology for sequencing purposes. The sequence was compared with the available data from National Centre for Biotechnology Information (NCBI) using Blast.

**RESULTS**

**Detection of cyanobacterial 16S rDNA**

Cyanobacterial 16S rDNA were amplified using primer CYA106F combined with CYA781R (a) and CYA781R (b) to generate a DNA fragment of 654 – 699 bp as shown in Figure 2.

All samples shown apparent bands on the gel electrophoresis image indicate the presence of cyanobacterial 16S rRNA (Table 3), showing the presence of the cyanobacteria in the samples.

**Detection of microcystin (mcyE) gene**

Generic microcystin (mcyE) gene were amplified using primer mcyE-F2 combined with mcyE-R4 to generate a DNA fragment of 809-812 bp as shown in Figure 3.

All samples shown positive 16S rRNA cyanobacterial gene indicates the presence of cyanobacteria in the samples.

**Detection of generic microcystin (mcyE) gene**

Microcystin (mcyE) gene was detected in the samples taken from Miri City Fan Lake while others samples shown negative results indicated that the absence of the microcystin gene.

**Detection of microcystin (mcyE) gene specific for *Microcystis* sp., *Anabaena* sp. and *Planktothrix* sp.**

Microcystin (mcyE) gene specific for *Microcystis* sp., *Anabaena* sp. and *Planktothrix* sp. were amplified using forward primer of mcyE-F2 combined with mcyE-R8 (*Microcystis* sp.), mcyE-12R (*Anabaena* sp.) and mcyE-plaR3 (*Planktothrix* sp.) to generate a DNA fragment of 247 bp as shown in Figure 4, Figure 5, and Figure 6 respectively.
PCR amplification for microcystin gene specific to Microcystis sp., Anabaena sp. and Planktothrix sp. shown no bands in gel electrophoresis images (Table 5), showing that microcystin gene was not amplified using the microcystin gene specific primers21.

Further PCR amplification were carried out to determine the genera producing the microcystin gene. However, detection of microcystin (mcyE) gene specific for Microcystis sp., Anabaena sp. and Planktothrix sp. showed negative results.

Following the negative results of the detection of microcystin (mcyE) gene specific for genera, the PCR products were cleaned and sent for sequencing to determine the genera producing microcystin gene. The sequence were compared with the available data from in National Centre for Biotechnology Information (NCBI) using BLAST.

**Table 1: Sequence of the primers used to amplify cyanobacterial 16S rRNA**

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequences (5’ to 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CYA106F</td>
<td>CGG ACG GGT GAG TAA CGC GTG A</td>
</tr>
<tr>
<td>CYA781R (a)</td>
<td>GAC TAC TGG GGT ATC TAA TCC CTT T</td>
</tr>
<tr>
<td>CYA781R (b)</td>
<td>GAC TAC AGG GGT ATC TCC CTT T TAA</td>
</tr>
</tbody>
</table>

*CYA781R primer is used as an equimolar mixture of CYA781R(a) and CYA781R(b).
* Two reverse primers were used as the both primers; CYA781R(a) and CYA781R(b) targets for morphologically different strains as the former targets filamentous cyanobacteria and the latter targets unicellular cyanobacteria13,14

Fig. 1: Map of sampling points in Miri, Sarawak. A) Taman Bulatan Lake, B) Miri City Fan Lake, C) Taman Hiltop Lake, D) Taman Awam Miri Lake, and E) Taman Tunku Lake.
Table 2: Sequence of the primers used to amplify mcyE gene

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequences (5’ to 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>mcyE-F2</td>
<td>GAA ATT TGT GTA GAA GGT GC</td>
</tr>
<tr>
<td>mcyE-R4</td>
<td>AAT TCT AAA GGC CAA AGA CG</td>
</tr>
<tr>
<td>mcyE-R8</td>
<td>CAA TGG GAG CAT AAC GAG</td>
</tr>
<tr>
<td>mcyE-12R</td>
<td>CAA TCT CGG TAT AGC GGC</td>
</tr>
<tr>
<td>mcyE-plaR3</td>
<td>CTC AAT CTG AGG ATA ACG AT</td>
</tr>
</tbody>
</table>

* Primer mcyE-F2 was used as forward primer for amplification of both generic and specific microcystin gene.

From the result of comparing the sequence with the available data in NCBI, the highest similarities of the sequence showed was the *Microcystis aeruginosa* (mcyE) gene 5'-CCCCGCCCCCTCCAGCTGAAATA

CCCGCCGGCTTCAATGCGGTATCAATGAAAC

ACCTTTTCTGCTTAAAACTGCGCACAAAGAC

GCTATCGGTACCGGTATGGTAGAAATTTAAGGAA

CGAAAGCTAATCAAGTAAAAGATAATGAACTT

TGACCCGACCCCTGTACATTTGTATATCATTTGAC

TGGCCGACGCTCTTTTTAAACGAGCCATTTGAC

CCTGCTCTTGCACCTACTCACAAATGATTTGC

ATACTGTTCACCTCTGCATATICTCAACGATCAGT

AACATTTTTTTTTTTTGTAGATGGAGCACAATTTA

AAATCTTTTTTTTTTTTGTAGATGGAGCACAATTTA

TGATTTCTCTCTAAAGATTTGATTCTTTGAAACATAG

CTTGGCTTTCTGTGACAAATTTGCGAAGGAAGCT

TCCG-3) resulted 66% identical to the available sequence in the GeneBank.

**DISCUSSION**

Both cyanobacterial 16S rRNA and microcystin genes were present throughout the study. Cyanobacterial 16S rRNA gene were detected in all samples collected in lakes located in Miri, Sarawak but microcystin (mcyE) gene were detected only in one out of twelve different field samples. In 2012, Harith and Hassan reported the presence of cyanobacteria in Ranchan Pool, Serian, Sarawak. 17 of cyanobacteria species was recorded from the
Table 3: PCR summary results for the presence of cyanobacterial 16S rRNA for the samples taken at different locations in Miri, Sarawak in June 2015. ‘+’ indicate the positive results while ‘–’ indicate the negative results.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taman Tunku</td>
<td>+</td>
</tr>
<tr>
<td>Taman Awam Miri 1</td>
<td>+</td>
</tr>
<tr>
<td>Taman Awam Miri 2</td>
<td>+</td>
</tr>
<tr>
<td>Miri City Fan 1</td>
<td>+</td>
</tr>
<tr>
<td>Miri City Fan 2</td>
<td>+</td>
</tr>
<tr>
<td>Miri City Fan 3</td>
<td>+</td>
</tr>
<tr>
<td>Taman Bulatan 1</td>
<td>+</td>
</tr>
<tr>
<td>Taman Bulatan 2</td>
<td>+</td>
</tr>
<tr>
<td>Taman Hilltop 1</td>
<td>+</td>
</tr>
<tr>
<td>Taman Hilltop 2</td>
<td>+</td>
</tr>
<tr>
<td>Taman Hilltop 3</td>
<td>+</td>
</tr>
<tr>
<td>Taman Hilltop 4</td>
<td>+</td>
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</tbody>
</table>

Fig. 3: Agarose gel electrophoresis image of 809-812 bp of PCR product obtained from the use of generic microcystin (mcyE) gene primers, mcyE-F2 as a forward primer combined with mcyE-R4 as a reverse primer on DNA extracts using GF-1 Bacterial DNA Extraction Kit (Vivantis) from samples taken at different lakes in Miri, Sarawak in June 2015. DNA was only visible for Miri City Fan 1, (Lane 7) while other samples show negative results. 1 = Ladder (VC 100bp); 2 = Empty; 3 = Negative control; 4 = Taman Tunku; 5 = Taman Awam Miri 1; 6 = Taman Awam Miri 2; 7 = Miri City Fan 1; 8 = Miri City Fan 2; 9 = Miri City Fan 3; 10 = Taman Bulatan 1; 11 = Taman Bulatan 2; 12 = Taman Hilltop 1; 13 = Taman Hilltop 2; 14 = Taman Hilltop 3; 15 = Taman Hilltop 4.
of cyanobacterial toxins should be alerted and monitored because the toxin production is gene specific.

Analytical methods are widely used for detection and quantification of cyanotoxins, but the half-life of the toxins are rather short and easily degraded in the environment. Thus, molecular techniques used to detect genes for toxin production in cyanobacteria in this study because the ability to produce the toxin is gene specific, not species specific. In the environment, microcystin is extremely stable in chemical hydrolysis and extreme high temperature (>300°C). Thus may accumulate in the extracellular from several days to years in the environment after toxic blooming occurrence. However, direct detection of toxins using method such as high performance liquid chromatography (HPLC) is difficult due to its vast different variants. In addition, microcystin is easily degraded by strong oxidation molecules such as ozone and breakdown by the aquatic bacteria such as Sphingomonas and Pseudomonas aeruginosa making toxin detection using analytical detection less reliable.

In detecting toxin-producing gene of Microcystis, Anabaena, and Planktothrix sp., researchers target mcyE gene from the mcyS gene cluster at the different primer targets. Presence of mcyE gene will immediately confirming the presence of toxic cyanobacteria in an environment as mcyE gene encode for protein important for the formation of 3-amino-9-methoxy-2,6,8-trimethyl-10-phenyl-4,6-decadienioic acid, (Adda) and glutamate-1-semialdehyde aminotransferase. Both molecules are important for toxicity. A research showed that...

Fig. 4: Agarose gel electrophoresis image of 247 bp of PCR product obtained from the use microcystin (mcyE) gene specific to Microcystis sp. primer, mcyE-F2 as a forward primer combined with mcyE-R8 as a reverse primer on DNA extracts using GF-1 Bacterial DNA Extraction Kit (Vivantis) from samples taken at different lakes in Miri, Sarawak in June 2015. Results were negative except for positive control (Lane 2). 1 = Ladder (VC 100bp); 2 = Positive control; 3 = Negative control; 4 = Miri City Fan 1

Fig. 5: Agarose gel electrophoresis image of 247 bp of PCR product obtained from the use microcystin (mcyE) gene specific to Anabaena sp. primer, mcyE-F2 as a forward primer combined with mcyE-12R as a reverse primer on DNA extracts using GF-1 Bacterial DNA Extraction Kit (Vivantis) from samples taken at different lakes in Miri, Sarawak in June 2015. Results were negative for the samples. 1 = Ladder (VC 100bp); 2 = Negative control; 3 = Miri City Fan 1
Fig. 6: Agarose gel electrophoresis image of 247 bp of PCR product obtained from the use of microcystin gene specific to *Planktothrix* sp. primer, mcyE-F2 as a forward primer combined with mcyE-12R as a reverse primer on DNA extracts using GF-1 Bacterial DNA Extraction Kit (Vivantis) from samples taken at different lakes in Miri, Sarawak in June 2015. Results were negative for the sample. 1 = Ladder (VC 100bp); 2 = Negative control; 3 = Miri City Fan 1

Table 5: PCR summary results for the presence of specific microcystin (*mcyE*) gene for the samples of Miri (City Fan) – Planktonic, which shown the positive result for generic microcystins. ‘+’ indicate the positive results while ‘–’ indicate the negative results

<table>
<thead>
<tr>
<th>Sample</th>
<th>Microcysts (<em>mcyE</em>) gene specific for <em>Microcystis</em> sp.</th>
<th>Microcysts (<em>mcyE</em>) gene specific for <em>Anabaena</em> sp.</th>
<th>Microcysts (<em>mcyE</em>) gene specific for <em>Planktothrix</em> sp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Miri City Fan 1</td>
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In conclusion, of twelve samples tested for microcystin encoding gene, only one sample was detected to possess the gene. The result shown the presence of potential risk of cyanobacterial toxin in one of Miri’s lake. This study revealed that this is the first time microcystin encoding gene was detected in Borneo Malaysia. The data presented was useful in water risk management and assessment of potential toxin contamination in recreational water in Sarawak.

ACKNOWLEDGMENTS

The authors are grateful to Universiti Sains Malaysia for providing financial assistance through USM RU grant and MOE ERGS grant.

Table 4: PCR summary results for the presence of generic microcystin (*mcyE*) gene for the samples taken at different locations in Miri, Sarawak in June 2015. ‘+’ indicate the positive results while ‘–’ indicate the negative results

<table>
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</tr>
<tr>
<td>Miri City Fan 3</td>
<td>-</td>
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<td>Taman Bulatan 1</td>
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<td>-</td>
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<tr>
<td>Taman Hilltop 1</td>
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<td>Taman Hilltop 2</td>
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<td>Taman Hilltop 3</td>
<td>-</td>
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<tr>
<td>Taman Hilltop 4</td>
<td>-</td>
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