Identification of Thermophilic Bacteria from Tirta Lebak Buana Hot Spring in Serang, Banten, Indonesia

Kenny Lischer

Bioprocess Engineering, Department of Chemical Engineering, Faculty of Engineering, Universitas Indonesia, Depok 16424, Indonesia, lischer.kenny@ui.ac.id

Follow this and additional works at: https://scholarhub.ui.ac.id/mjt

Recommended Citation
DOI: 10.7454/mst.v25i3.3993
Available at: https://scholarhub.ui.ac.id/mjt/vol25/iss3/6

This Article is brought to you for free and open access by the Universitas Indonesia at UI Scholars Hub. It has been accepted for inclusion in Makara Journal of Technology by an authorized editor of UI Scholars Hub.
Identification of Thermophilic Bacteria from Tirta Lebak Buana Hot Spring in Serang, Banten, Indonesia

Kenny Lischer*

Bioprocess Engineering, Department of Chemical Engineering, Faculty of Engineering, Universitas Indonesia, Depok 16424, Indonesia

*E-mail: lischer.kenny@ui.ac.id

Abstract

Since Taq polymerase was first explored and identified from thermophilic bacteria, these bacteria have become well-known sources of thermostable enzymes. New thermophilic bacteria have been investigated to broaden biodiversity and translation research. Studies have shown interests in Indonesia because of thermophilic bacteria found in hot springs. This country is traversed by the ring of fire and has more than 70 volcanoes, resulting in the wide distribution of hot springs across the country. Although many reports have been performed, studies have yet to explore thermophilic bacteria in Tirta Lebak Buana hot springs, Java Island, Indonesia. This research was the first to examine thermophilic bacteria in Tirta Lebak Buana hot spring. Two samples from two different sampling sites were obtained and analyzed through 16srRNA analysis (sampling sites A and B). Measurements indicated that the temperature (50 °C) in sampling site A was higher than that in sampling site B (40 °C), but they had similar pH (7.0). Polymerase chain reaction (PCR) showed that the 16srRNA of the specimen was around 1465 bp. The analysis of the 16srRNA sequence revealed that the obtained bacteria have a similar sequence and close relationship with Bacillus subtilis subsp. stercoris strain N12.

Keywords: hot spring, moderate thermophile, thermophilic bacteria, tirta lebak buana, 16srRNA

1. Introduction

Initially discovered in 1953, thermophilic bacteria can remain alive at high temperatures (more than 40 °C–90 °C) [1]. Since the exploration of Taq polymerase from Thermus aquaticus, thermophilic bacteria have led to the global revolution of biology [2–5]. They have become well-known microorganisms for their ability to produce thermostable enzymes that can be used for industrial applications [6]. However, the biodiversity of thermophilic bacteria has not yet been fully explored.

Indonesia has more than 70 volcanoes distributed in Sumatra, Java, Bali, Kalimantan, Sulawesi, and Papua [7], and a total of 256 hot springs are found in Indonesia [8]. Interestingly, hot springs are known as the habitat of thermophilic bacteria [1]. Therefore, thermophilic bacteria from hot springs in Indonesia are interesting...
organisms to be explored. Microbes in hot springs around Indonesia were initially explored in 1991. Approximately eight different species were identified from nine selected hot springs, namely, Toya Bungkah, Sikidang, Candradimuka, Sileri, Domas, Badak, Djarian, Ciater, and Gunung Gede, respectively [9]. Since then, efforts devoted to biodiversity exploration in hot springs have significantly increased.

At least three methods, namely, culture and morphological approach, 16srRNA, and whole-genome sequencing, can be applied to investigate microbial diversity in hot springs. Culture and morphological approaches are considered the easiest methods to identify microbes. These methods usually involve strain staining and morphological characterization [10–13]. However, these methods are limited by the small number of specimens to be identified [14]. With whole-genome sequencing, bacteria can be identified in detail from a cellular level to a DNA level. This method can also be utilized to explore all functional genes and improve the chance to determining newly discovered species [14]. Although the cost of whole-genome sequencing has decreased, it is still considerably high in developing countries. Another identification technique at the DNA level is 16srRNA analysis. 16srRNA is a gene that can be used for bacterial fingerprinting [15,16]. This method is considerably cheap because PCR is the only technique performed to target and sequence a single gene. As a result, 16srRNA becomes a commonly used method for the exploration of thermophilic bacteria, as demonstrated in hot springs in Prataan (Paenibacillus sp.), Cisolok (Paenibacillus cisolokensis), and Tutung (Brevibacillus borstelensis and Paenobacillus sp.) [17–19].

Indonesia has numerous microbiologically explored hot springs [9, 18, 20, 21]. Although extensive research has been conducted, studies have yet to be performed to explore thermophiles in Tirta Lebak Buana hot springs. Therefore, this research aimed to isolate and identify thermophilic bacteria in Tirta Lebak Buana through 16srRNA analysis.

2. Methods

Sample collection and characterization. Water from two sites (A and B) in Tirta Lebak Buana Hotsprings in Serang, Banten (6°32’55.2” S 106°23’58.0” E) was collected in vacuum-insulated water bottles in October 2019. It was then analyzed in terms of temperature, pH, and sulfur content. The samples were transported at room temperature and stored at -20 °C. They were cultured in plates containing 1.6% peptone, 0.8% yeast extract, 0.6% sodium chloride, 2 g/L MgSO₄, and 1.6% gellan gum (Thermofischer, USA) and incubated at 50 °C and 40 °C.

Species identification. The obtained colonies were cultured in a Luria-Bertani medium and incubated at 37 °C. Then, genomic DNA was extracted by using a Quick-DNATM Fungal/Bacterial Miniprep kit (Zymo Research, D6005, USA). Polymerase chain reaction (PCR) was performed to the target 16srRNA by using the following forward and reverse primers, respectively: 5’-AGAGTTTGATCTGGCTCAG-3’) and 5’-CGGTTACCTGGTACGA CTT-3’). During PCR, Taq polymerase and buffer (Bioneer, Korea) were also used at 92 °C for 5 min, 60 °C for 0.5 min, and 72 °C for 1 min for 32 cycles followed by a final elongation at 72 °C for 5 min [22, 23]. The PCR results were analyzed through gel electrophoresis.

Phylogenetic tree and multiple alignment sequence analysis. Bidirectional sequencing was conducted to identify the 16srRNA sequence. The obtained sequenced was then used to analyze phylogenetic trees via BLAST. Multiple alignment sequence comparison with several known bacteria was performed with Clustal Omega.

3. Result and Discussion

Sampling location. Numerous hot springs are distributed in various locations in Indonesia. One of them is Tirta Lebak Buana hot spring, where thermophilic bacteria have yet to be explored. In Figure 1a, Tirta Lebak Buana hot spring is located in Banten province, Java Island, Indonesia (6°32’55.2” S and 106°23’58.0” E). The location is less than 100 km from Krakatau. Water was collected from two different sampling sites for bacterial isolation. The main pond where the heat of hot spring is generated was set as sampling site A (Figure 1b). The part of the hot spring where people take a bath was assigned as sampling site B (Figure 1b). Both hot springs have neutral pH and no sign of sulfur odor (Table 1). They differ in temperature, i.e., the temperature in site A (50 °C) is higher than that in site B (40 °C). This temperature indicates that thermophilic bacteria may be thermophiles or moderately thermophiles.

PCR of 16srRNA. The thermophilic bacteria that can be obtained from this hot spring are unknown. In this study, water obtained from both sampling sites was cultured. The results revealed white colonies in the culture (Table 1). The obtained culture was then analyzed by using 16srRNA methods. In Figure 2, the size of the PCR band from sites A and B was around 1465 bp. This result showed that the 16srRNA of the thermophilic bacterial cultures was successfully amplified. The size of the band was considerably in the correct position because of the forward primer 27F and the reverse primer 1492R; this result indicated that it could amplify 16srRNA from sequence number 27 to 1492 [22, 23].
16srRNA sequence analysis. The obtained 16srRNA was sequence from the PCR bands from sites A and B to gather further information about the species. The sequence was analyzed through BLAST application to find information regarding closely related bacteria. The result revealed that the samples from sites A and B had the closest relationship with *Bacillus subtilis* subsp. *stercoris* strain N12 (Figure 3). Multiple alignment sequencing was performed to further confirm the relationship of obtained bacteria with *B. subtilis* subsp. strain N12. In Figure 4, the samples from sites A and B have identical 16srRNA sequence. They also have a similar 16srRNA sequence to that of *B. subtilis* subsp. strain N12 (99% similarity). Neither the sequence of *Escherichia coli* nor the sequence of *Geobacillus thermoleovorans* is similar to both samples. Therefore, both samples may be *B. subtilis* subsp. *stercoris* strain N12. This strain is characterized as moderate thermophilic bacteria or facultative thermophilic bacteria because they can live actively at high temperatures but below 40 °C [24]. However, moderate thermophiles have optimal growth temperature between 60 °C and 80 °C [25]. The only discovery that has been made is its identification in a food-waste bioreactor [24]. Interestingly, it can also be found in nature, specifically in Tirta Lebak Buana hot spring. Future studies should be performed to identify the bacterial strain in detail via whole-genome sequencing. Various moderate thermophiles have many thermostable enzymes, such as SpCas9, lipase, cellulase, protease, and esterase [26–28]. Therefore, thermostable enzymes from *B. subtilis* subsp. *stercoris* strain N12 should be elucidated in future research.

**Table 1. Sampling Site Characteristics**

<table>
<thead>
<tr>
<th>Sampling location</th>
<th>T (°C)</th>
<th>pH</th>
<th>Sulfur</th>
<th>Colony Color</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site A</td>
<td>50</td>
<td>7.0</td>
<td>-</td>
<td>White</td>
</tr>
<tr>
<td>Site O</td>
<td>40</td>
<td>7.0</td>
<td>-</td>
<td>White</td>
</tr>
</tbody>
</table>

![Figure 1. Sampling Location. (a) Map Location of Tirta Lebak Buana Hot Spring. (b) Sampling Sites A and B](image1)

![Figure 2. PCR Amplification of 16srRNA of the Samples from Sites A and B](image2)

![Figure 3. Phylogenetic Trees of Samples from Sites A and B](image3)
4. Conclusions

This study is the first to isolate thermophilic bacteria from Tirta Lebak Buana hot spring. The obtained bacterial culture may be a moderate thermophile, *B. subtilis* subsp. *stercoris* strain N12. This promising result can be used for further studies related to its morphology, biological significance, and thermostable enzymes for industrial purposes. This study provides opportunities for future research, especially on *B. subtilis* subsp. *stercoris* strain N12, which should be further explored.

*Makara J. Technol.*
Acknowledgments

KL performed the research, analyzed the data, and wrote the article. We would like to thank the Department of Chemical Engineering for the publication support.

References