Gene Encoding Nitrilase from Soil Sample of Lombok Gold Mine Industry using Metagenomics Approach

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Abstract

This paper describes an efficient screening gene nitrilase from contaminated soil from Lombok gold mine industry. DNA was extracted directly from soil using the soil DNA isolation kit based on enzymatic, chemical and mechanical lysis. The existence of nitrilase gene in soil sample can be identified by nitrilase gene amplification using H1F-H1R primer. BLASTN analysis result revealed that the nitrilase gene fragment which was amplified by H1F-H1R primer has a high homology with *Rhodococcus rhodochrous* strain *tg1-A6 nitrilase gene*. These amplification and DNA fragment sequencing results indicated that nitrilase gene existence on soil sample can be identified by metagenomic approach.

Keywords: Metagenomics; nitrilase gene; *Rhodococcus rhodochrous*; soil.

1. Introduction

The biosphere is dominated by microorganisms however mostly they have not been studied. Traditional methods for culturing microorganisms limit analysis to those that grow under laboratory condition [1]. Demand for novel biocatalysts is continuously increased and thereby prompting the development of novel experimental approaches to find and identify novel biocatalyst-encoding genes. Recently, there has been an increase in the number of studies using a metagenomics approach to investigate the catalytic potential of uncultured microorganisms [2].

In the search for novel biocatalysts, there are various metagenomic strategies used for discovery of enzymes for a broad spectrum of biotechnological applications. Enzymes are key to all biosynthetic pathways in the production of natural metabolites in microbial system. Direct cloning has led to important discoveries of genes responsible for novel biological functions. New genetic information on various industrial enzyme such as metagenomic approaches [3].

Nitrilases are enzymes that hydrolyse non peptide carbon-nitrogen bonds. They are important industrial enzymes used to convert nitriles (R-C≡N) such as cyanide, directly into corresponding acids and ammonia. Nitrilases are quite rare in bacterial genomes.
and less than 20 were reported before the application of metagenomics for their detection in environmental DNA. Two studies targeting environmental genomes report the detection of more than 337 novel nitrilases [4]. This has dramatically increased of interest on studying nitrilases and the newly discovered diversity can be applied for the enantioselective production hydroxyl carboxylic acid derivates [5, 6]. The variety of newly discovered nitrilases represents a broad diversity of nitrilase sequence space.

Nitrilases have been promoted to be a valuable alternative to chemical hydrolysis in an organic chemical processes because of their mild reaction conditions and reduced output of environmental pollution [7]. However, environmental problems caused by cyanide are due to the production of large amounts of cyanide-containing wastes by human industrial activities, such as production of organic nitriles, nylon and acrylic plastics, paints, drugs and the use of cyanide in gold mining and the metal industries [8].

Gold mining in developing countries can be organized and regulated with mechanisms in place to protect workers and the environment. But mining can also informal and small-scale with minimal regulatory control of activities. This second form of mining is described as artisanal or small-scale gold mining (ASGM). Recently, the increased gold price can be seen to the extent of operations to the island of Lombok and Sumbawa. Physicochemical extreme environment such as highly polluted environments contain a low microbial community size. Microbe that is capable of living in this hostile environment has evolved special mechanisms for survival. Due to the low community size and biomass of these habitats are not as readily available as other environments by metagenomic approaches [9]. For this purpose, an efficient screening gene nitrilase from contaminated soil from gold mining by metagenomes approach have been introduced.

2. Materials and Method

2.1. Research Location

The research was conducted through a field survey of the Sekotong (South West Lombok), Keruak and Sekarbela during Mei 2011. Cyanidation and mercury tailings were sampled directly from tailings dams. On Lombok, hard-rock mining occurs throughout the Sekotong region, with ore transported as far as capital city Mataram for processing (Sekarbela). But in Sumbawa, amalgamation and cyanidation facilities have been constructed close to the point for ore extraction. The extent of mining activity is increasing in both Sekotong and Sumbawa. These samples had been chosen due to suspected
elevated metal content, and a number of cyanide and nitrile degrading bacteria had previously been isolated from these soil.

2.2. Soil preparations

2.2.1. DNA extraction

Soil DNA extraction was performed using the soil DNA isolation kit (Mobio laboratories, USA) based on enzymatic, chemical and mechanical lysis.

2.2.2. DNA quantitation

For quantitating DNA, quantitation of double stranded DNA was calculated from the optical density absorbance value measured by a spectrophotometer (ABI) using wavelengths of 260 nm and 280 nm. PCR condition. Reaction mixture for PCR contained: 25 μL go tag buffer 2.5 μL complete, 0.5 μL dNTPs, 1.0 μL primer 1, 1.0 μL primer 2, 0.5 μL (1 unit) DNA Taq polymerase, 1 μL DNA, 1.5 μL MgCl₂ 25 mM, 18 μL ddH₂O in final volume 50 μL. DNA amplification was performed with thermocycler Takara (Japan) by using the followed program: 10 min hotstart at 94°C followed by 35 cycles consisting of denaturation (2 min at 94°C), annealing (60 s at 52°C), extension (90 s at 72°C) and a final extension step at 72°C for 5 min. PCR product was analysed in 1% agarose gel electrophoresis run with TAE buffer (40 mM Tris-acetate pH 8.1 mM EDTA) and stained with ethidium bromide. The primer used for nitrilase encoding gen amplification are listed in Table 1, Genomic DNA of *Rhodococcus pyridinovorans* TPIK was used as positive control.

2.2.3. Gel electrophoresis

After PCR amplification, the amplified PCR products were checked for expected size on 1.5% (w/v) agarose gel (LE analytical grade Promega, Madison, USA). Agarose gel was loaded with 10 μL of each PCR amplified product and run in 1 x TAE. A ready-to-use 1Kb plus DNA Ladder-molecular weight standard (Novagen, USA) and negative control were run together with the PCR amplified products. The PCR product was separated by electrophoresis system at a constant voltage of 80 V for 50 min. Then, the gel was stained with ethidium bromide staining (0.5 μg · mL⁻¹) for 30 min and followed by washing with distilled water for about 3 min. Finally, the gel was visualized under UV transilluminator, and photos were taken using gel documentation system.

2.2.4. Gel extraction (Qiagen)

DNA fragment was excised from the agarose gel with a sharp scalpel and weigh the gel slice in a colorless tube. The gel was suspended in buffer QG (three volume buffer QG to one volume gel) and incubated at 50°C for 10 min. Every 2 min to 3 min solution was mix well to help dissolve gel. After the gel slice has dissolved completely add one
gel volume of isopropanol and mixed gently upside down. The mixture was applied to the quick column then centrifuged for 1 min. The flow-through was discarded and placed column back in the same collection. 750 μL of Buffer PE was added to quick column and centrifuged for 1 min. The flow-through was discarded and centrifuged the column for additional then placed column into a clean 1.5 mL microcentrifuge tube. DNA was elute with Buffer EB or TE buffer and stored at −20°C.

2.2.5. Nucleotide sequencing and sequence analysis

Nucleotide sequences were determined via 1st BASE (Singapore) and analysed using blastn or blastx software \cite{10} from the GeneBank (NCBI) database.

3. Results and Discussions

3.1. DNA purity

The purity of the DNA extracts is important for gene amplification by PCR and further analysis such as DGGE fingerprinting or cloning of amplicons. To examine the purity of DNA extracts, the UV absorptions at 260 nm and 280 nm were measured and the ratio of 260 : 280 were calculated. The quality of the extracted DNA was evaluated by the $A_{260}$ : $A_{280}$ ratio and values high than 1.8 indicate a good DNA extract without protein or humic acid contamination \cite{11}. The DNA samples have OD $A_{260}$ : $A_{280}$ ratio around 0.613 to 1.3. DNA extracts of the soil sample showed absorbance ratio below 1.8, which reflects the co-extraction of contaminant. Soil is particularly complex matrix containing many substances, such as humic acids, which can be co-extracted during DNA isolation. Removal of humic acids is essential before the DNA can be processed further. Because of this, a range of DNA purification techniques has been developing. The resulting DNA extracts were analysed by agarose gel electrophoresis (see Figure 1). Although DNA extract of the sample showed the below the detection limit, but showed sharp band in gel electrophoresis.

Soil is a particularly complex matrix containing many substances, such as humic acids, which can be co-extracted during DNA isolation. Removal of humic acids is essential before the DNA can be processed further. Because of this, a range of DNA purification techniques has been developed. Sephadex G-200 spin columns have proven to be one of the best to remove contaminants from soil DNA. Recently, a pulse field electrophoresis procedure using a two-phase agarose gel, with one phase containing polyvinylpyrrolidone (PVPP), was developed for removal of humics \cite{12}.

Critical steps in direct soil DNA extraction methods are lysis of microbial cells and eliminating humic acids, protein, and other inhibitors. For cell lysis to be effective, mechanical treatment should be followed rather than chemical ones. Various physical and chemical treatments have been evaluated for cell rupture, which include shaking the sample in lysis buffers, detergents or glass beads, inclusion of lysozyme \cite{13}.

The ultra clean soil kit DNA isolation kit remains the most highly cited purification method for soil in use today. Inhibitors like Humic acid, pulvic acids, polysaccharides
Figure 1: Electrophoresis of DNA extracted from soil samples, 6 \( \mu \)L of eluted DNA was loaded on 1% agarose gel, and electrophoresis for 55 min Lane 1 = 1 Kb DNA ladder Marker; Lane 2, DNA extract.

and polyphenols are removed using IRT (Inhibitor Removal Technology) for the highest success rates in PCR of any commercial method. The ultraclean DNA isolation is fully compatible with bead beating technology if harder beating for lysis of tough organism is desired and provided the highest quality DNA based on successful amplification functional genes of nitrilase. We generated the expected 900 bp to 1100 bp fragment with soil extract from mining gold in Sekotong, Mataram.

3.2. PCR detection of nitrilase genes

Following PCR using DNA extract from soil as the template, the amplification product was detected using agarose gel electrophoresis. As a shown in Figure 2 a specific DNA fragment (about 900 bp), which matched the correct size of the nitrilase gene was successfully amplified. Nevertheless, only smears of non-specific fragment were obtained indicating that may the primer were cross reacted with other DNA sequence.

The gel was extracted by QIAquick gel extraction kit for remove the non-specific fragment, the removed slice of gel should contain the desired DNA inside. *Nitrilase gene* fragment sequencing results of the soil samples were obtained base sequence of 957 bp. Those sequenced have been compared with another gene sequence of domain bacteria using the BLASTN program at NCBI webpage. Furthermore analysis indicated that sequence are closely related to *Rhodococcus rhodochrous* strain tgl-A6 *nitrilase gene* with 99% homology.
4. Conclusion

The existence of nitrilase gene in soil sample from Lombok gold mine industry can be identified by direct DNA extraction and nitrilase gene amplification using H1F-H1R primer. BLASTN analysis result revealed that the nitrilase gene fragment which was amplified by H1F-H1R primer has a high homology with Rhodococcus rhodochrous strain tg1-A6 nitrilase gene.

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