Sequential Isolation of Saturated, Aromatic, Resinic and Asphaltic Fractions Degrading Bacteria from Oil Contaminated Soil in South Sumatera

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Abstract

Sequential isolation has been conducted to obtain isolates of saturated, aromatic, resin, and asphaltene fractions degrading bacteria from oil contaminated sites. Five soil samples were collected from South Sumatera. These were analyzed using soil extract medium enriched with oil recovery or Remaining-Oil recovery Degradated (ROD) as sole carbon and energy sources according to the isolation stage. ROD at the end of every isolation stage analyzed oil fractions by use of the SARA analysis method. Six isolates of bacteria have been selected, one isolate was fraction saturates degrading bacteria that are *Mycobacterium sp.* T1H2D4-7 at degradation rate 0.0199 mgs/h with density 8.4x10⁶ cfu/g from stage I. The isolate T2H1D2-4, identified as *Pseudomonas sp.* was fraction aromatics degrading bacteria at accelerate 0.0141 mgs/h with density 5.1x10⁶ cfu/g are obtained at stage II. Two isolates namely *Micrococcus sp.* T3H2D4-2 and *Pseudomonas sp.* T1H1D5-5 were fraction resins degrading bacteria by accelerate 0.0088 mgs/h at density 5.6x10⁶ cfu/g and 0.0089 mgs/h at density 5.7x10⁶ cfu/g are obtained at stage III. Isolation of stage IV has been obtained two isolates *Pseudomonas sp.* T4H1D3-1and *Pseudomonas sp.* T4H3D5-4 were fraction asphaltenes degrading bacteria by accelerate 0.0057 mgs/h at density 5.6x10⁶ cfu/g and accelerate 0.0058 mgs/h at density 5.7x10⁶ cfu/g.

Abstrak

Isolasi Bertahap Bakteri Pendegradasi Fraksi Jenuh, Aromatik, Resin, dan Aspal dari Tanah Terkontaminasi Minyak di Sumatra Selatan. Penelitian isolasi bertahap telah dilakukan untuk mendapatkan bakteri pendegradasi fraksi jenuh, aromatik, resin dan aspal. Isolasi dilakukan terhadap lima sampel tanah terkontaminasi minyak dari Sumatera Selatan. Medium isolasi menggunakan Soil extract diperkaya oil recovery atau oil recovery sisa degradasi (OSD) sebagai satu-satunya sumber karbon dan energi sesuai tahapan isolasi. OSD setiap akhir tahap isolasi difraksinasi menggunakan SARA Analysis untuk mengetahui fraksi jenuh, aromatik, resin dan aspal. Hasil penelitian mendapatkan enam isolat bakteri terpilih berdasarkan kecepatan degradasi tertinggi pada setiap tahap, satu isolat bakteri pendegradasi fraksi jenuh yaitu Mycobacterium sp. T1H2D4-7 dengan laju degradasi 0,0199 mg/jam dan kepadatan 8.4x10⁶ cfu/gram dari tahap I. Isolat T2H1D2-4 teridentifikasi sebagai *Pseudomonas sp.* merupakan bakteri pendegradasi fraksi aromatik dengan laju degradasi 0,0141 mg/jam dan kepadatan 5,1x10⁶ cfu/gram diperoleh pada tahap II. Dua isolat yaitu Micrococcus sp. T3H2D4-2 dan Pseudomonas sp. T1H1D5-5 merupakan bakteri pendegradasi fraksi resin yang masing-masing mempunyai laju degradasi 0.0088 mg/jam dengan kepadatan 5.6×10^6 cfu/gram, dan 0,0089 mg/jam dengan kepadatan $5,7x10^6$ cfu/gram diperoleh dari tahap III. Isolasi tahap IV diperoleh dua isolat yaitu Pseudomonas sp. T4H1D3-1 dan Pseudomonas sp. T4H3D5-4 yang merupakan bakteri pendegradasi fraksi aspal, masing-masing mempunyai kecepatan degradasi 0,0057 mg/jam dengan kerapatan 5,6x10⁶ cfu/gram, dan 0,0058 mg/jam dengan kerapatan 5.7×10^6 cfu/gram.

Keywords: aromatics, asphaltenes, hydrocarbon-degrading bacteria, saturates, resins, sequential isolation

1. Introduction

Environmental pollution with petroleum and petroleum products has been recognized as one of the most serious

current problems especially when associated with accidental spills on a large-scale [1]. Likewise, soil contamination by oil is common in oil and gas industry activities [2]. Ueno *et al.* [3], explained that oil

contamination in soil was caused by, among others, the occurrence of oil spills or spills during transport, pipeline leaks, tank-cleaning of crude oil storage tanks. and also because of accidents like oil well blow-outs. Petroleum contaminated soil, can affect the biota either above or below ground level. These impacts can lead to death (lethal effects), physical damage to biota (subeffect), and habitat degradation due to lethal contamination of a-biotic factors on the soil. In this regard, the decree from the Minister of Environment No. 128 of 2003 [4] required the processing of petroleum-contaminated soil in an attempt towards environmental restoration, through bioremediation. The use of bioremediation to change the characteristic and composition of the oil contaminants to become harmless is by reducing the mobility, mass, and concentration of petroleum contaminants in the soil.

Generally the fractions of petroleum hydrocarbons are saturates, aromatics, resins, and asphaltenes fractions [5-6]. The content of the four fractions in oilcontaminated soil is determined by the length of time oil is exposed to the soil. Fractions of petroleum hydrocarbons that are volatile (saturates and aromatics fractions by atomic C<8) evaporate in the environment. Thus the fraction of hydrocarbon-contaminated soil contained on the relative has a greater molecular weight and the level of biodegradability is more difficult [7]. The process of biodegradation of petroleum hydrocarbons is influenced by several factors, the composition and concentration of hydrocarbon petrophilic (hydrocarboncompounds, bacteria degrading bacteria), environmental factors during the process of biodegradation, and necessary nutrients [8]. To perform bioremediation of oil-contaminated soil, the mixed bacterial cultures in the form of a consortium have potential application in the bioremediation of the polluted site by removing all the fractions of hydrocarbon pollutants from the environment [9].

Some research on the isolation of hydrocarbondegrading bacteria have been done (Sutiknowati, 2011; Sivaraman et al., 2011; Mandri and Lin, 2007; Mittal and Singh, 2009), but they only obtained bacteria that can degrade specific hydrocarbons fractions, but cannot all the fractions. Fraction degrade aromatic (naphthalene, phenantrene, trichlorodibenzofuran, and benzo[a]pyrene) degrading bacteria isolated from "beach simulator tank" was added with nutrients N and P [10]. Pseudomonas mendocina was isolated from bilge oil contaminated water at Mormugao harbor. This saturated strain effectively degraded fractions (tetradecane, hexadecane and octadecane) leaving a residual concentration of about 73%, 54% and 40% respectively in 120 hours [11]. Three bacterial isolates have been obtained from soil contaminated with diesel and engine oils, they were identified as Flavobacterium sp., Acinetobacterium calcoaceticum and Pseudomonas

aeruginosa. All isolates were capable of degrading saturates fraction (*n-paraffin*) up to 80% in a 2 week period [12]. *Pseudomonas* strain PS-I has been isolated from soils contaminated with crude oil spill, this isolate could degrade saturates (*alkanes*) (70.69%) and aromatics (45.37%) fractions [13]. Therefore, sequential isolation needs to isolate the bacterial strains that can degrade all fractions of hydrocarbons petroleum (saturates, aromatics, resins, and asphaltenes fractions).

The area of South Sumaterais one area associated with the oil industry, both the public and private sectors. One side effect of oil industry activity is contamination of soil by oil. Sequential isolation is the isolation method based on a succession of bacterial communities that process biodegradation of hydrocarbon petroleum contaminants. Hopefully, through sequential isolation, we can obtain petrophilic bacteria capable of using the carbon from all fractions of petroleum components. Subsequently this may serve as the basis for the development of biological agents in the bioremediation of oil-contaminated soil.

2. Methods

Samples of oil-contaminated soil was collected from five locations: Abab (D1), Benakat (D2), Limau (D3), Raja (D4), and Talangjimar (D5).Contaminated soils at each sampling site and map of sampling sites are seen in Figure 1. Five locations in sequence have been contaminated by petroleum for five, four, three, two, and one year respectively. Sampling was conducted using multiple sampling methods, by determining the five sites based on stratified sampling, while the determination of the three sub-locations of the depth of the surface (0 cm), middle (15 cm), and bottom (30 cm) at each location with a stratified sampling. From each sampling point 0.5 to 2.0 kg of contaminated soil was taken using a soil sampling tool (auger), samples from the same depth were pooled and labeled, then inserted into the sample container aseptically [8,14].

Twenty-five grams of contaminated soil was put into 500 mL erlenmeyer flash containing 225 mL Soil Extract Medium (SEM) made from the same soil sample and 15% (w/v) of oil recovery was added as the sole carbon and energy source. Cultures were incubated at room temperature (26-31 °C) in an orbital shaker with 100 rpm for one week. Every 48 hours a sample was taken and diluted using 1 mL physiological saline solution (0.85% NaCl) up to 10^{-8} , respectively, then each dilution of 0.1 mL was taken and inoculated into the medium Atlas Oil surfaces Agar (AOA) using the spread method. AOA medium composition was prepared according to the Atlas [15] g per 990 mL: to 15; KH₂PO₄ 1.0, K₂HPO₄ 1.0, NH₄NO₃1.0, 0.2 MgSO₄.7H₂O; FeCl₃ 0.05; CaCl₂.7H₂O 0.02; and oil 10 mL. Each culture was incubated at room temperature



Figure 1. Map of Sampling Sites of Petroleum-Contaminated Soil (left), Oil-Contaminated Soils at Each Sampling Site (Right)

(26-31 °C) for 3 x 24 hours, and then colonies having different morphological features were purified by the quadrant streaking method. The Remaining-Oil recovery Degradated (ROD) was sterilized and reused in stage II as the sole carbon source. Isolation of stage II was performed in the same way as isolating stage I but using the ROD stage I as the sole carbon source, and so on until reaching the isolation of stage IV (Figure 2). Oil Recovery and the ROD of each stage of isolation analyzed saturates, aromatics, resins, and asphaltenes fractions using the SARA Analysis method (Figure 3).

Bacterial isolates obtained at each stage of isolation were verified to determine the best ability of the isolates in degrading the oil fractions. Each bacterial isolate was inoculated 2.5% (v/v) cell density of 2.5 $\times 10^8$ cfu/mL into a 100 mL glass bottle containing 12.5 mL of Mineral Medium (MM) with crude oil, a composition according to the Atlas [15] grams per Liter: K₂HPO₄ 4.5, (NH₄)₂SO₄ 1.0, MgSO₄.7H₂O 0.2, NaCl 0.1, CaCl₂ 1.0, FeCl₃ 0.02 and enriched with 250 mg oil recovery as the sole carbon and energy source, and one glass bottle was not inoculated to function as the control. Cultures were incubated at room temperature (26-31 °C) at 100 rpm in an orbital shaker for six days, this experiment used completely randomized designs with 3 times replications for each isolate. Furthermore, cell numbers were counted using the spread method, and analysis of the fraction of saturates, aromatics, resins, and asphaltenes using the SARA Analysis method as shown in Figure 3.

Fraction analysis was performed after the cultures were taken prior to the analysis of cell numbers. 20 mL npentane was then added to two hundred and fifty to five hundred milligrams of oil recovery and ROD 1 up to ROD 4 in the culture bottles, homogenized and separated from the medium by a separator funnel. The asphaltenes fractions was obtained by filtering the fraction of n-pentane using a filter paper of known weight. Furthermore, the remaining filtrate collected



Figure 2. Typical Scheme for Isolation of Hydrocarbon-Degrading Bacteria by Sequential Isolation Method



Figure 3. SARA Analysis Method for Separating Oil into Saturate, Aromatic, Resin and Asphaltene (SARA) Components

from the separation of the asphaltenes is commonly known as maltenes. It contains the remaining three fractions, saturates, aromatics and resins. These three fractions are separated using open- chromatographic column with silica gel (60-100 mesh). Saturates fraction on percolation in n-pentane eluant, are not absorbed on activated silica under the conditions specified. The arom saturates fraction of the oil was eluted from column with 20 mL of n-pentane at 5 mL/min. The solvent was removed by evaporation to recover the saturates fraction. Aromatics fraction are absorbed by activated silica in the presence of n-pentane, and desorbed by saturates of the saturates with a saturates of the saturates

fraction. Aromatics fraction are absorbed by activated silica in the presence of n-pentane, and desorbed by toluene after removal of the saturates under the conditions specified. The aromatics fraction of the oil is eluted from the chromatographic column using 20 mL toluene at 5 mL/min. The resin fraction of the oil is eluted from the chromatographic column using 20 mL 90:10 toluen : methanol solution at 5 mL/min. All fractions were determined by the gravimetry method and then reported as milligrams [16-17]. The degradation rate of the fraction is determined using the following equation:

The degradation rate (mg/h) =
$$\frac{KF0-KFt}{\Delta t}$$
 (1)

 KF_0 where is the initial concentration of fraction, KF_t is the concentration fraction after *t* time, Δt is the difference. The data of the rate of degradation fraction of saturates, aromatics, resins, and asphaltenes was analyzed by F test to determine the isolates with the highest activity to degrade each fraction tested by the New Duncan's Mutiple Range Test (DNMRT) at significant level (α) <5%.

Identification and characterization of selected isolates were performed by morphological colonies on agar stab, agar slant, agar plate using temperature growth, Gram and endospores staining, motility, and biochemical [1,18-21]. Results obtained were used for further identification on the basis of Bergey's Manual of Determinative Bacteriology [22] and Bergey's Manual of Systematic Bacteriology [23].

3. Results and Discussion

The results of the sequential isolation from five locations were grouped according to the stages of isolation. We obtained 14 isolates of bacteria at stage I, all of these isolates are degrading the fraction of saturates. In stage II, we obtained two saturates degrading isolates and 15 aromatics degrading isolates. Isolation of stage III obtained 14 resins degrading isolates, and more isolation results are presented in Table 1. Thus there is indication of the existence of a trend pattern between the stages of isolation with the degraded fraction. In stage I, all isolates tend to degrade saturates; isolates from stage II tend to degrade the

aromatics fraction. Isolates obtained in stage III tend to degrade the resins fractions, and isolates from stage IV tend to degrade the asphaltenes fraction. The trend occurs because each group of isolates differ in starting work to degrade each fraction, the group of saturates degrading bacteria starts first, then is followed by aromatics degrading bacteria, resins degrading bacteria, and asphaltenes degrading bacteria, respectively. It is also supported by concentrations of saturates, aromatics, resins, and asphalt fractions at each stage isolation (Figure 4). Figure 4 shows that instage I ROD-1 the saturated fraction decreases dramatically, while the other factions experience no real decline. Respectively it shows that at stage II ROD-2, stage III ROD-3, and stage IV ROD-4 the aromatics, resins, and asphaltenes fractions decrease dramatically. This condition explains that the fraction of saturates starts to degrade rapidly at stage I so bacterial isolates obtained at this stage are saturates fraction degrading isolates. Likewise, aromatics, resins, and asphaltenes fractions degraded starting at stage II, III, and IV respectively, so that at stage II the isolates of bacteria degrading are aromatics fractions, at stage III and IV bacterial isolates obtained is for resins degrading bacteria and asphaltenes degrading bacteria respectively. These results are in accordance with the theory put forward by Taki et al. [7] that the order of ease of biodegradation of petroleum hydrocarbon fractions from the easiest to the most difficult is the fraction of saturates, aromatics resins, and asphaltenes.

Table 1. Number of Isolates base on Isolation Stage

Stage	Oil fractions						
	saturat.	aromatics	resins	asphalt.			
Ι	14	-	-	-			
II	2	15		-			
III	-	-	14	-			
IV	-	-	-	11			



Figure 4. Average Concentration of Saturates, Aromatics, Resins, and Asphalthenes Fractions in the Initial Condition and at Each Stage of Isolation

Result of verification test indicated that there were 16 saturates degrading isolates, the rate of each saturates degrading isolates are listed in Figure 5. The highest rate was the isolate T1H2D4-7 from Raja, this isolate has the rate of degradation of the aromatics fraction of 0.0199 mgs/h on the number of cells 8.4×10^6 cfu/g.

As 15 isolates were obtained from aromatic fractions, all isolates from stage II and the rate of each aromaticsdegrading isolates are listed in Figure 5. The isolateT2H1D2-4 was the only one isolate that has the highest degradation rate of aromatic fraction, from Benakat. This isolate has the rate of degradation of the

aromatic fraction of 0.0141 mgs/h on the number of cells 5.1×10^6 cfu/g.

Figure 6 shows the resins fraction degrading isolates from stage III, from 13 isolates of resin degrading isolates there are two with the highest rate of degradation of isolate T3H2D4-2 and T1H1D5-5 from Raja and Talangjimar respectively. The degradation rate of each is 0.0088 mgs/h at cell number 5.6×10^6 cfu/g and 0.0089 mgs/h on the number of cells 5.7×10^6 cfu/g respectively.



Figure 5. Degradation Rate of Saturates Degrading Isolates (Left), Degradation Rate of Aromatic Degrading Isolates (Right), the Stem Given the Same Lower Case Notation Indicates Non-Significant Difference at the α <0.05



Figure 6. Degradation Rate of Resin Degrading Isolates (Left), Degradation Rate of Asphalthene Degrading Isolates (Right), the Stem Given the Same Lower Case Notation Indicates Non-Significant Difference at the α <0.05



Figure 7. Cell Morphologies of Selected Isolates, (a) *Mycobacterium sp.* (T1H2D4-7), (b) *Pseudomonas sp.* (T1H1D5-5), (c) *Pseudomonas sp.* (T2H1D2-4), (d) *Micrococcus sp.* (T3H2D4-2), (e) *Pseudomonas sp.* (T3H3D5-1), and (f) *Pseudomonas sp.* (T4H3D5-4)

Characteristics	Code of Isolates					
	T1H2D4-7	T2H1D2-4	T1H1D5-5	T3H2D4-2	T4H1D3-1	T4H3D5-4
Morpho. colony						
- NA stab	Echinulate	Echinulate	Echinulate	Echinulate	Echinulate	Echinulate
- NA plate						
Configurations	Circular	Irregular	Circular	Rhizoid	Circular	Circular
Margins	Entire	Laborate	Entire	Filiform	Entire	Undulate
Elevations	Convex	Raised	Convex	Hilly	Convex	Convex
Structures	Smooth	Concentric	Smooth	Contoured	Smooth	Smooth
Colours	White	Yellow	Yellow	Yellow	Yellow	Orange
- NA slant	Filiform	Effuse	Spreading	Rhizoid	Filiform	Echinulate
- NA 37 °C	Aerob	Aerob obligat	Aerob	Facultative	Aerob obligat	Aerob obligat
- NA 41 °C	Aerob	Aerob obligat	Aerob	Facultative	Aerob obligat	Aerob obligat
Morphol. Cell						
- Gram	Positive	Negative	Negative	Positive	Negative	Negative
- Cell shape	Rod	Rod	Rod	Coccus	Rod	Rod
- Endospores	No spores	No spores	No spores	No spores	No spores	No spores
Motility	Non motil	Motil		Non motil	Motil	Motil
Biochem. test						
- Sitrat	Negative	Negative	Negative	Positive	Negative	Negative
- H ₂ S	Negative	Negative	Negative	Negative	Negative	Negative
- MR	Negative	Negative	Negative	Positive	Negative	Negative
- VP	Negative	Negative	Negative	Negative	Negative	Negative
- Hyd. Gelatin	Positive	Negative	Negative	Positive	Positive	Negative
- Urease	Negative	Positive	Positive	Negative	Positive	Positive
- Indole	Negative	Negative	Negative	Negative	Negative	Negative
- Hyd. Starch	Positive	Negative	Negative	Negative	Negative	Negative
- Fer. Glucose	Positive	Negative	Positive	Positive	Positive	Positive
- Fer. Lactose	Positive	Positive	Positive	Positive	Positive	Positive
- Fer. Sucrose	Positive	Negative	Negative	Positive	Negative	Positive
- Catalase	Positive	Positive	Positive	Negative	Positive	Positive
- Hyd. Fat	Positive	Positive	Positive	Negative	Negative	Positive
- Hyd. Casein	Negative	Negative	Negative	Negative	Negative	Negative
Degrader of fraction	Saturates	Aromatics	Resins	Resins	Asphaltenes	Asphaltenes
Genera	Mycobacterium sp	Pseudomonas sp	Pseudomonas sp	Micrococcus sp	Pseudomonas sp	Pseudomonas sp
	(T1H2D4-7)	(T2H1D2-4)	(T1H1D5-5)	(T3H2D4-2)	(T4H1D3-1)	(T4H3D5-4)

Table 2. The Results of the Characterization and Identification of Six Selected Bacterial Isolates

The group of asphaltenes fractions degrading isolates are 11, all isolates are from stage IV (Figure 6). Isolates that have the highest rate are isolates T4H1D3-1 namely0.0057 mgs/h at number of cells 5.6×10^6 cfu/g, this isolate is from Limau and T4H3D5-4 namely 0.0058 mgs/h at number of cells 5.7×10^6 cfu/g, this isolate is from Talangjimar.

Characterization and identification of bacterial isolates were performed on six selections based on the highest rate in fractions degradation. Six bacterial isolates included one isolate of saturates degrading bacteria, one isolate of aromatics degrading bacteria, two isolates of resins degrading bacteria, and two isolate of asphaltenes degrading bacteria. The cell morphology of six isolates can be seen at Figure 7, and then more results of the characterization and identification are listed in Table 2.

4. Conclusion

The sequential isolation method is applicable to obtain petrophilic bacteria that can degrade saturates, aromatics, resins, and asphaltenes fractions. Six bacteria were sequentially isolated from oil-contaminated soil in South Sumatra, one isolate of saturates-degrading bacteria of *Mycobacterium sp.* (T1H2D4-7) from Raja, with degradation rate 0.0189 mgs/h with the number of cells 8.4×10^6 cfu/mL. One isolate identified as *Pseudomonas sp.* (T2H1D2-4) from Benakat was aromatics degrading bacteria with a rate of 0.0141 mgs/h with the number of cells 5.1×10^6 cfu/mL. Two isolates namely *Micrococcus sp.* (T3H2D4-2) from Raja and *Pseudomonas sp.* (T1H1D5-5) from Talangjimar were resins degrading with a rate of 0.0088 mgs/h on cell numbers 5.6×10^6 cfu/mL and 0.0089 mgs/h on the

number of cells 5.7×10^6 cfu/mL respectively. Two isolates namely *Pseudomonas sp.* (T4H1D3-1) from Limau and *Pseudomonas sp.* (T4H3D5-4) from Talangjimar are asphaltenes degrading with a rate of 0.0057 mgs/h on the number of cells 5.6×10^6 cfu/mL and 0.0058 mgs/h on the number of cells 5.7×10^6 cfu/mL respectively.

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