Anaerobic degradation of crude oil by sulphate reducing bacteria isolated from soils contaminated with petroleum hydrocarbons

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Abstract

This study included isolation a mix cultures of sulphate reducing bacteria from soils in Shaeba regain contaminated with petroleum hydrocarbons by using API medium enriched with oxygen reducing agents and saturated with gas (90% N₂) and (10% CO₂) by using sodium lactate as sole carbon source. The result showed that the number of bacteria ranged (4 x 10⁵ – 4.5 x 10⁵) cells / g of soil. Also using crude oil as a sole of carbon source to study the ability of sulphate reducing bacteria on degradation of crude oil under anaerobic condition. The result showed these bacteria have high ability for degradation of crude oil. The percent rate of degradation in culture was (84.4 %) after (54) days incubation and the rate concentration of total petroleum hydrocarbons (TPH) extraction from culture value (22.06) μg / L, while in control was (190.84) μg / L.

Introduction

The bane of industrial progress has been the generation and release into the environment of the huge amounts of toxic compounds which have caused widespread contamination of the land and water. These chemical have been released into the environment creating countless number of contaminated sites, most widely distributed environmental pollution can be attributed to hydrocarbons contamination, caused by oil tanker accidents, storage tank rupture, transport accidents, and old petrol station (Jain et al, 2005 and Aoshima et al., 2006).

Crude oil is a complex mixture of hydrocarbons, basically composed of aliphatic, aromatic and asphaltene fraction along with nitrogen, sulfur and oxygen contain compounds (Aoshima et al., 2006). Some of these compounds have been reports to carcinogenic, mutagenic and have immunodulatory effects on humans, animal and plant life (Miller & Maller, 1981 and Van-Gesel et al., 2001). Removed of the hydrocarbons from contaminated environments involved physical and chemical processes which expensive and some time difficult to execute, so the microbial
The degradation of spilled hydrocarbons is a major technique in natural decontamination processes, converting toxic organic to harmless products, often carbon dioxide and water, also its can degrade a wide range of hydrocarbons (Saadoun, 2002 and Koma et al., 2003). The most processes of hydrocarbons biodegradation occurs in aerobic condition but there are several indicating the anaerobic degradation occurs in the absent of oxygen by Sulphate Reducing Bacteria (Balk, 2007 and Widdel et al., 2007).

The sulphate – reducing bacteria (SRB) are large group of anaerobic organisms that play important role in many biogeochemical processes.(Barton and Hamilton, 2007). The main property of this group is obligate anaerobic bacteria population is their active use of sulphate as a final electron acceptor during anaerobic respiratory and capable of generation hydrogen sulphide (H2S) from the reduction of sulphate (Boetius et al.,2000 and Sahrani et al., 2008).

SRB utilize very wide spectrum of different low molecular compounds (Lactate, acetate, propionate, succinate, format, pyruvate, ethanol and aliphatic acid) as carbon and energy sources (Caumette, 1993). Sulphate and organic matter concentration, temperature and salinity are the main environmental factors controlling the number and distribution of SRB and the rate of bacterial sulphate reduction (Mudryk et al., 2000). They are widely distributed in nature from anaerobic mud found at the bottom of the ocean to the intestines of humans. (Gad and White, 1996). Also were found to grow environmental contaminants such as petroleum hydrocarbon constituents (benzene, toluene, ethylbenzene, xylenes and alkanes), petroleum reservoirs and oil production facilities (Zhang and Young, 1997 and Barton and Hamilton, 2007). SRB using to purify industrial wastewater from heavy metals and in recent years studies have also been made on the use of SRB for the biodegradation of organic matter (Sanir et al., 2001; Kleikemper et al., 2002 and Rezscycka et al., 2004).

Hence, the aim of this study was isolated and numerated of SRB from contaminated soil and study their ability for anaerobic degradation of crude oil.

Material and methods
Samples collection
Soil samples were collected from Shaeba reign near the south oil refinery company in Basrah which is highly contaminated with crude oil, the samples collected from depth soil at (15-30) cm under surface by clean handle, especially in sterile plastic containers and sealed to avoid oxygen contamination and under semi anaerobic condition and transferred immediately to the laboratory (Rooney-Varga et al., 1997).

Isolation and cultivation of sulfate reducing bacteria
SRB were isolated from soil samples by using liquid API medium API, (1975) which is used as selective growth medium, with the following compositions: yeast extract (1 g); MgSO4. 7H2O (0.2 g); Fe(NH4)2(SO4)2. 6H2O (0.2 g); NaCl (1 g); K2HPO4 (0.01 g); Ascorbic acid (0.1 g) and Sodium lactate (2.24 g). All of the dry chemical were measured out first and prepared by adding to each a liter of distilled water, the pH was adjusted to (7.2) using (1 M) NaOH solution, medium sterilized by autoclave.
under (121°C) and pressure (15) bound / inch² for 15 minute, left to cool at room temperature. This medium treated with added oxygen reducing agents has the following: Sodium dithionate (0.3 g / l) and L-systein (0.28 g / l) these compounds were sterilized by heat, and saturated under gas phase (90% N₂) with (10% CO₂) before being inoculated with the samples (Rabus et al., 1996 and Teske et al., 1996).

(Figure1). The soiled medium was prepared by added (1.5 % wt / vol) agar. To isolate of SRB from samples (1 g) of sample mixed with (9 ml) of liquid API medium and shuck well until it become homogenized, (1ml) of homogenized mat surface layer was transfer to screw tubes containing approximately full of liquid API medium under condition mentioned in isolation, tubes were sealed by screw cover and coated with paraffin tape to prevent diffusion of O₂ into medium, and incubated at (35 °C) until blackening of the medium was recorded as positive for SRB presence. (Hirnes et al., 1999 and Carignan et al., 1994).

Enumerated of SRB

For determination of cell numbers, dilution series (10⁻² - 10⁻⁹) was prepared from each samples with liquid API medium under condition identical to that mention in isolation of SRB in screw stopped tubes. From (10⁻⁵-10⁻⁷) dilutions, (0.5) ml was incubated into triplicate anaerobic roll tubes containing approximately full of API agar medium at (45) °C, tubes were stopped well and coated with paraffin tape, then tubes were rolling between two hand to speared inoculums through the medium, incubated at (35) °C until well formed colonies become visible (Hines et al., 1999), numbers of cells measured as following:

Numbers of cells / g of soil = Numbers of colonies × 1/dilution.

Purification of SRB

The colonies obtained from cultivation series dilution in API agar were picked by means of finely drawn sterile Pasteur pipettes, the colonies were immediately transferred into tube of fresh liquid medium, community of SRB formed by mixing equal volumes of eight cultures of bacterial communities originating from various samples (Rabus et al., 1996).

Biodegradation of crude oil by SRB

Using north rumella crude oil in the chemical experiment to determined ability of SRB for degraded crude oil, stored in cool place by using sterilized container until being used. Conical flasks (100) ml with (80) ml of sterilized liquid API medium without sodium lactate, added (3%) of crude oil by sterilized pipettes to every conical, then inoculums by (3) ml of activated mix cultures of SRB, incubated at (35) °C for (54) days in dark without shaking at form semi sloping to maximum the contact area between oil and medium, conical flasks were sealed and coated with paraffin tape, control sample was prepare by added (3%) crude oil to medium only and incubate under same conditions (Ruetter et al., 1994 and Rabus et al., 1996).

Extraction of oil hydrocarbons

The oil consumption ratio by SRB in liquid medium and control samples, estimated by the weight measurement method, extraction (80) ml growth culture with (100) ml of carbon tetrachloride in separation funnel, was well
mixing many times, the mixture of solvent and culture was leave to separate to form two layers, the lower layer (contain oil hydrocarbons) was collected and transferred into separation Column contain an anhydrous sodium sulfate which make up absorbed water and other contaminates, hydrocarbons fraction was collected in round bottle, and volatilized of solvent by rotary evaporator under low pressure at (55) °C, no increase than (60) °C. The weight of oil extraction was measured after drying from solvent (UNEP, 1992), oil consumption ratio was calculated from residual oil components as the following:

\[
\text{Degradation rate\%} = \frac{\text{mg of crude oil control} - \text{mg of crude oil test}}{\text{mg of crude oil control}} \times 100
\]

Total Petroleum Hydrocarbons (TPH) was determined in testing and control samples, dissolved dry hydrocarbon extraction with n-hexane (5) ml, then were measured by spectrofluorometer system (type Shimadzu- RF540) was equipped with recording (type Shimadzu - DR3), emission at (360) nm and Excitation at (310) nm. The control sample was measured under same condition, results compared with standards curve as shown in figure (5) for north rumela crude oil by making serial dilution.

**Results**

The bacterial cultures showed good growth anaerobically in API medium and on agar medium that supplied with oxygen reducing agents and presence of sodium lactate as carbon source under (90% N2) with (10% CO2). Rapid growth was observed on this medium as it only took about three days for the liquid medium and one to two days on agar medium to turn blackening due to sulfide production, the observation of black color might imply the present of SRB (figure 2). Black colonies characteristics as single isolates were examined on API agar appear (figure 3). SRB were enumerated in (1g) of soil samples in solid API medium with about (4 x 10^-5 - 4.5 x 10^-5) cells/g of soil.

The results showed high capacity of SRB for growth in presence of crude oil as sole carbon source, the initially clear medium become black, because of bacterial growth and production of iron sulphide (FeS) after sulphate reducing and emulsification of oil, camper with control samples without SRB present (figure 4). Degradation ability percentage for SRB was (88%) after incubation (54 days). Mean total petroleum hydrocarbons (TPH) remaining in cultivated culture for SRB was (22.06 µg/L) while the control was (190.84 µg/L) (Table 1).

**Discussion**

The cultures of SRB by use liquid API medium with sodium lactate as carbon source and oxygen reducing agents under (90% N2) with (10% CO2) condition, showed good growth within three days, because most SRB are growth on lactate. The growth was estimated on the basic of the amount of sulphate reduced in the medium to turn blackening. (Hanselmann et al., 1995 and Rzeczycka et al., 2004)
The results of SRB numeration was showed large numbers of these bacteria, this result is referred to ability of SRB for growth in environments contaminated with oil hydrocarbons, and utilized of compounds which found in crude oil which contain amount of \( \text{SO}_4^{2-} \) necessary for growth as final electron accepter, this is a good agreement with the findings Rabus et al. (1996) and Zhang and Young (1997), also during the degradation of crude oil, low molecular weight organic acids such as acetate, propionate and butyrate which in turn may serve as carbon sources for SRB (Cozzarelli et al., 1994). Moreover, numerous studies have shown that SRB genera are known to readily degrade a wide range of organic acids and associated with the degradation of the respective carbon sources in many environments (Widdle and Bak, 1992; Hanselmann et al., 1995; Purdy et al., 1997; Sass et al., 1998 and Kuever et al., 2001). SRB number in the soil range between \( (4 \times 10^5 \text{ to } 4.5 \times 10^5) \) cell per g wet soil, this number highest to the extent that is predominated in hydrocarbon contaminated soil, because of optimum environment for these microbial (Machaughton et al., 1999).

In the present study growth of SRB on crude oil as sole sources of carbon showed value have been degraded in which that \( (88.4\%) \) of its through ability to growth and turned black, compared with control samples, this result is agreed with (Ruerer et al., 1994 and Rabus et al., 1996), in which SRB utilize aliphatic and aromatic hydrocarbons directly from oil samples as the only source of organic substance under anoxic condition. The ability for growth wide variety of SRB in crude oil due to their contain many sources of reducing equivalent for sulphate reduction and organic carbon for cell synthesis (Kleikemper et al., 2002).

SRB represents a special enzymic system such as pyrophosphate, ATP sulphylase, bisulphite reductase and desulphofucin (Mudryk et al., 2000), therefore these enzymes increase their ability to hydrocarbons degradation and toleration to high rate of contaminants (Safinowski et al., 2004 and widdle et al., 2007).

Figures and table:

(Figure 1) Saturated media with \text{N}_2 \text{ and CO}_2

(Figure 2) Growth of SRB in API medium
(Figure 3) Isolated colonies of SRB in solid API medium

(Figure 4) A: Degradation of crude oil by SRB

\[ Y = -4.77303 + 960111(x) \]
\[ r = 0.994 \]

(Figure 5) Standard curve for Rumella crude oil
(Table 1) Degradation percentage and main Total Petroleum Hydrocarbons (TPH)

<table>
<thead>
<tr>
<th>SAMPLES</th>
<th>DEGRADATION %</th>
<th>MAIN TPH µG/L</th>
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</thead>
<tbody>
<tr>
<td>SRB cultures</td>
<td>84.4</td>
<td>22.06</td>
</tr>
<tr>
<td>Control</td>
<td></td>
<td>190.84</td>
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Reference


التكسير اللاهوائي للنفط الخام بوساطة الجراثيم المختزلة للكبريت المعزولة من الترب الملوثة

بالهيدروكاربونات النفطية

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الخلاصة

تضمنت الدراسة الحالية عزل مزارع خليطية من البكتيريا المختزلة للكبريت من الترب الملوثة بالنفط الخام من منطقة الشعبية جنوب العراق باستخدام وسط API المدعوم بالعوامل المختزلة للأوكسجين والمشتتات بما في ذلك N₂ (90 %) و (CO₂ 10 %) و (N₂ + CO₂) (10%) (x 4.440 x 10⁻⁷ ملم / غم من التربة. كما تم استخدام مزيج النفط الخام كمصادر وحيض للكربون في دراسة قابلية البكتيريا المختزلة للكبريت على تكييف النفط الخام تحت ظروف لاهوائية، وأظهرت النتائج وجود قابلية عالية للبكتيريا على تكييف النفط الخام، إذ بلغ معدل نسبة الملوحة للتكسير 84.4 % في مزارع الجراثيم بعد (54) يوماً من الحضن، مما بلغ معدل تركيز الهيدروكاربونات النفطية الكلية المستخلص من مزارع الجراثيم (2006 ميكروغرام / لتر أما في عيدات السيطرة فقد بلغ (190.84 ميكروغرام / لتر. 