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Evaluation of bacterial diversity recovered from petroleum samples using different physical matrices



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ABSTRACT

Unraveling the microbial diversity and its complexity in petroleum reservoir environments has been a challenge throughout the years. Despite the techniques developed in order to improve methodologies involving DNA extraction from crude oil, microbial enrichments using different culture conditions can be applied as a way to increase the recovery of DNA from environments with low cellular density for further microbiological analyses. This work aimed at the evaluation of different matrices (arenite, shale and polyurethane foam) as support materials for microbial growth and biofilm formation in enrichments using a biodegraded petroleum sample as inoculum in sulfate reducing condition. Subsequent microbial diversity characterization was carried out using Scanning Electronic Microscopy (SEM), Denaturing Gradient Gel Electrophoresis (DGGE) and 16S rRNA gene libraries in order to compare the microbial biomass yield, DNA recovery efficiency and diversity among the enrichments. The DNA from microbial communities in petroleum enrichments was purified according to a protocol established in this work and used for 16S rRNA amplification with bacterial generic primers. The PCR products were cloned, and positive clones were screened by Amplified Ribosomal DNA Restriction Analysis (ARDRA). Sequencing and phylogenetic analyses revealed that the bacterial community was mostly represented by members of the genera *Petrotoga*, *Bacillus*, *Pseudomonas*, *Geobacillus* and *Rahnella*. The use of different support materials in the enrichments yielded an increase in microbial biomass and biofilm formation, indicating that these materials may be employed for efficient biomass recovery from petroleum reservoir samples. Nonetheless, the most diverse microbiota were recovered from the biodegraded petroleum sample using polyurethane foam cubes as support material.

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Introduction

Crude oil biodegradation in petroleum reservoirs affects the world production of fuels, making the recovery and refining processes more expensive. For many years, the prevalent occurrence of biodegradation in petroleum wells has been attributed to the aerobic bacterial degradation of hydrocarbons, which can be stimulated by oxygen carried by the infiltration of meteoric waters in the reservoir.¹ However, there is strong evidence for the widespread occurrence of obligate anaerobes in subsurface petroleum systems,^{2–4} and the flushing of meteoric water does not indicate that highly reactive oxygen survives transportation to deep reservoirs, since even small concentrations of organic compounds can remove oxygen from an aquifer.

In recent work, researchers have suggested that biodegradation processes can occur at the oil–water transition zone, in which microbial life should be possible within water droplets containing active microbial communities.⁵ Data gathered from several studies indicate that oil biodegradation in deep subsurface petroleum reservoirs occurs through anaerobic microbial metabolism rather than aerobic mechanisms, resulting in a decrease of light hydrocarbons and an increase of oil density, acidity, viscosity and sulfur content.^{6–8} In addition, viable anaerobic hydrocarbon degradation processes have recently been established for both saturated and aromatic hydrocarbons.^{9–11} Studies aiming to evaluate intermediate metabolites, characteristic of anaerobic hydrocarbon degradation, have been carried out and allowed the identification of compounds such as reduced 2-naphthoic acids,¹² 2-methylnaphthalene, tetralin, as well as naphthoic acids in petroleum reservoirs.¹³

Sulfate reduction and methanogenesis are the most likely processes responsible for in-reservoir hydrocarbon oxidation.¹⁴ Oil degradation linked to sulfate reduction would explain the consistent hydrocarbon compositional patterns seen in many degraded oils worldwide. Sulfate arises from geological sources, such as evaporitic sediments and limestone, or from the injection of seawater for pressure stabilization, and may lead to significant oil degradation and increased residual-oil sulfur content.¹⁵ Souring in oilfield systems is most commonly due to the action of sulfate-reducing prokaryotes, a diverse group of anaerobic microorganisms that respire sulfate and produce sulfide (the key souring agent) while oxidizing diverse electron donors.⁸

In this sense, efforts have been made by several researchers in order to recover and characterize the anaerobic microbial community inhabiting the deep petroleum biosphere.^{2,16,17} The study of genomes of uncultivated microbes have become possible through metagenomics, a cultivation-independent approach that allows to explore the metabolic potential of the unseen biodiversity by cloning large DNA fragments directly isolated from the environment.¹⁸ With the use of the metagenomic approach, bacteria capable of degrading petroleum hydrocarbons, including anaerobes, have been more deeply investigated and their metabolic routes unraveled.^{19,20} However, the extremely low amount of DNA obtained from samples derived from petroleum reservoirs using direct nucleic acid extraction procedures is often a restraint when the

phylogenetic and/or metabolic diversity of microbial communities are investigated,^{21,22} because of the low cellular density and activity found in such hostile environment. Microbial enrichments using different culture conditions, simulating the chemical and physical parameters of natural environments, can be applied to overcome this limitation and increase the recovery of DNA from environments with low cellular density.²³ Although cultivation under laboratory conditions can diminish the biodiversity recovered, this technique allows the selection of microorganisms that have some function of particular interest, such as enzymatic activity or biodegradation ability.²⁴

This work aimed to evaluate the efficiency of different matrices, used as physical supports, in recovering anaerobic bacterial diversity from a biodegraded oil sample derived from a petroleum reservoir in Campos Basin (Brazil). The matrices were used in order to evaluate their effect in the increase of biomass, as well as a support for biofilm formation. The relative abundance and diversity of the anaerobic microbiota recovered from the enrichments were compared by using microscopic and molecular analysis (DGGE and 16S rRNA libraries).

Material and methods

Sampling

Petroleum samples were obtained in July 2005 from five production wells at the Pampo Platform, Campos Basin (Macaé, RJ, Brazil), with logistic support from CENPES/Petrobras, as described in details by Vasconcellos et al.²² Samples were collected in triplicate using 500 mL sterilized Schott bottles, which were completely filled with the samples in order to avoid oxygen influx. Samples were kept on ice during transportation to the laboratory and stored at room temperature for subsequent anaerobic bacterial enrichment assays.

Anaerobic enrichments

The biodegraded petroleum sample (P2) used in this work as inoculum (10% v/v) for the anaerobic enrichments was collected from oil reservoir 2 in Campos Basin, RJ, Brazil.²² This well was characterized as highly biodegraded, level 5–6, according to Peters and Moldowan,²⁵ with average temperature 71 °C and approximately 2000 m deep. The petroleum sample was homogenized in water bath at 50 °C. The enrichments were settled in Schott bottles (1 L) containing 500 mL of Zinder medium²⁶ supplemented with organic substrates (sodium acetate, sodium formate, sodium lactate, yeast extract, methanol) to stimulate the growth of sulfate reducing bacteria, according to methods described by Dubourgier et al.²⁷ and Silva et al.²⁸ The cysteine–HCl solution (2 mM) was added to the enrichments (1%, v/v) as final electron acceptor.

Three different matrices were independently applied as physical supports to allow bacterial biofilm formation and increase biomass recovery under sulfate reducing condition: (1) polyurethane foam cubes (PF) (1 cm²), (2) slices of shale

(S), and (3) arenite (A). Bacterial enrichment using P2 as inoculum but without any matrix was settled as control (WS). Each condition was evaluated in triplicate.

The polyurethane foam cubes were submitted to UV sterilization for 20 min followed by immersion in autoclaved Zinder medium, under anaerobic condition (N_2 flow). Approximately 80 cubes were added to each flask of microbial enrichment under condition (1). Samples of shale and arenite, which are natural components of oil reservoirs, were kindly donated by Dr. Eugenio V. dos Santos Neto (CENPES/Petrobras). Shale was sliced in small pieces, while arenite was grated in powder. Both materials were autoclaved twice in Schott bottles (250 mL) containing 200 g of slices or powder. After sterilization, 10 g of each matrix were inoculated in the corresponding microbial enrichments.

All anaerobic enrichments were incubated at 55 °C, during 60 days, in a rotary shaker at 100 rpm.

Microscopic analysis

Scanning electronic microscopy (SEM) was used to evaluate the abundance and diversity of cell morphology of all bacterial enrichments under study. The analyses were developed with a Zeiss microscope model LEO 982, at Embrapa/CNPMA, using protocols described by Melo et al.²⁹.

DNA extraction

The DNA extraction from the bacterial enrichments was carried out using a protocol based on Großkopf et al.³⁰ and Neria-González et al.,³¹ with adaptations to petroleum samples. Firstly, 50 mL of a sterilized Tween 80 solution (10%) (Sigma–Aldrich) were added into enrichments in order to promote homogenization of oil/water phases, as well as improving the recovery of cells adhered on the physical supports (foams, shale and arenite). Total enrichment volume was distributed in sterile tubes (50 mL) and centrifuged at 10,000 rpm, for 20 min, 4 °C. Supernatants were discarded and cells were transferred to microtubes (2 mL). Afterwards, microbial pellets retrieved from the enrichments (3 × 500 mL) were suspended in 600 µL PBS buffer, homogenized by vortex and lysozyme was added at a final concentration of 17 mg/mL. After incubation at 37 °C for 2 h, proteinase K and SDS were added (final concentration of 0.7 mg/mL and 2%, respectively) and the solution was incubated at 60 °C for 90 min. The microtubes were submitted to three freeze–thaw cycles (2 min in liquid nitrogen followed by 2 min at 65 °C). Glass beads were added to the tubes and manual agitation was performed for 1 min. The solution was extracted once with equal volume of saturated phenol (pH 8.0) and once with equal volume of chloroform:isoamyl alcohol (24:1). For DNA precipitation, 5 M NaCl (10%) and 2 volumes of cold ethanol were added to the solution. The pellet was washed once with ethanol 70%, dried and suspended in Milli-Q water. The yield and integrity of the DNA obtained were confirmed through NanoVue Plus™ Spectrophotometer (GE Healthcare) and electrophoresis in 1% agarose gel stained with ethidium bromide and documented using a UVP BioImaging System GDS-8000 (UVP, Upland, CA, USA).

Construction of 16S rRNA gene libraries, ARDRA and phylogenetic analyses

For the construction of the 16S rRNA gene libraries, amplification was performed from total community DNA, obtained from each enrichment, by using the bacterial primer set 27f and 1100r.³² Only one library was assembled for each enrichment. Fifty microliter-reaction mixtures were made contained 50 ng of total DNA, 2 U of Taq DNA polymerase (Invitrogen), 0.2 mM of dNTP mix and 0.4 µM of each primer, in 1X Taq buffer. The PCR amplifications were performed using 10 cycles of 1 min at 94 °C, 30 s at 60 °C, decreasing 0.5 °C each cycle, and 3 min at 72 °C, followed by another 10 cycles of 1 min at 94 °C, 30 s at 56 °C and 3 min at 72 °C. Amplicons were pooled from five reactions (~500 ng), purified using GFX™ PCR-DNA and gel band purification kit (GE Healthcare) and cloned using the pGEM-T cloning vector kit, according to the manufacturer's instructions (Promega, Madison, Wisc.). Insert-containing clones were submitted to ARDRA by digestion of M13 amplicons with the enzymes Hae III, Hha I and Msp I, independently, at 37 °C for 2.5 h. Clones representing distinct ribotypes were selected for DNA sequencing and phylogenetic affiliation.

The 16S rRNA gene sequences were determined by direct amplification of the inserts from overnight grown clone cultures with M13 forward and reverse primers and sequencing with the DYEnamic ET Dye Terminator Cycle Sequencing Kit for the automated MegaBace 500 system (GE Healthcare) using the primers 10f, 1100r, 765f and 782r,³² according to the manufacturer's recommendations. Partial 16S rRNA gene sequences obtained from clones were assembled in a contiguous sequence using the phred/Phrap/CONSED program.^{33,34} Phylogenetic affiliation was achieved as described previously by Vasconcellos et al.²².

The nucleotide sequences determined in this study were deposited at the Genbank database under the accession numbers: GenBank ID: JN998802 to JN998890.

DGGE analyses

The PCR targeting 16S rDNA for the DGGE analyses was performed using the universal primers 968f (attached to a 40-nucleotide GC-rich sequence) and 1401r,³⁵ which are homologous to the conserved bacterial 16S rDNA regions. The PCR amplifications were performed in 50 µL reactions containing 50 ng of total community DNA recovered from the microbial enrichments, 5 µL of 10× Tris–HCl reaction buffer, 1.5 mM MgCl₂, 0.4 µM primers 968f and 1401r, 0.2 mM dNTP mix and 2 U Taq DNA Polymerase (Invitrogen, Grand Island, N.Y., USA). The PCR amplifications were performed using an initial denaturation step of 5 min at 94 °C, 10 cycles of 1 min at 94 °C, 30 s at 58 °C, decreasing 1 °C each cycle, and 2 min at 72 °C, followed by another 25 cycles of 1 min at 94 °C, 30 s at 53 °C and 2 min at 72 °C. The amplicons were first checked on 1.2% agarose gels prior to the DGGE analyses.

The DGGE analyses were carried out in the D-Code Universal Mutation Detection System (Bio-Rad, USA) using a linear denaturing gradient of urea and formamide ranging from 50% to 70% (100% denaturant corresponding to 7 M urea and 40% (v/v) deionized formamide). Gels (6% polyacrylamide)

Table 1 – Amount of DNA extracted from anaerobic enrichments using different supports and without support.

Anaerobic enrichment	DNA amount (ng/ μ L)
Shale	15 \pm 1.52
Arenite	14 \pm 0.57
Polyurethane foams	25 \pm 2.64
Without support	20 \pm 2.44

containing 6 μ L of PCR products for each sample, in triplicate, were run at 50 V and 60 °C for 14 h in 0.5 \times TAE buffer. Gels were stained with SYBR Green 1 \times solution and documented under UV light.

Results

Microscopic analyses of bacterial enrichments

A dense cellular biomass (up to 10⁸ cells/mL) was observed when using polyurethane foams as matrices in the anaerobic enrichments after 60 days of incubation. The enrichments without physical supports exhibited low medium turbidity (10⁴ cells/mL) when compared to the others in which matrices were employed.

The SEM analyses demonstrated that bacterial enrichments without physical supports yielded low abundance of cells and no biofilm formation (Fig. 1a and b). Actually, in this condition cells were shown to be sparsely distributed. On the other hand, a dense biomass yield and biofilm formation could be observed around (shale) or inside the porous (arenite and polyurethane foam) of the other matrices (Fig. 1c–h).

A predominance of coco rods was detected in the enrichments without physical supports (Fig. 1a and b), whereas straight rods forming polymeric structures (EPS) were observed when using polyurethane foams (Fig. 1c and d). The use of arenite and shale as supports allowed an intense biofilm formation involved by EPS, and thus the determination of the microbial morphology was not possible (Fig. 1e–h).

DNA extraction and DGGE analyses

After 60 days of incubation, the recovery of community DNA was possible for all the petroleum-based anaerobic enrichments performed in this study. It is worth to mention that different controls were set up, using only the matrices and sterilized medium. The controls did not show any turbidity, demonstrating that no contamination occurred. Although the enrichment without physical support exhibited lower turbidity of cells, the enrichments containing shale and arenite as support showed lower DNA recovery when compared to the enrichments without support and to the one containing polyurethane foams (Table 1). This was probably due to the strong adsorption of bacterial cells to the pieces of shale and arenite granules, making it difficult the recovery of all the biomass developed in these enrichments.

The DGGE analyses revealed distinct band profiles between samples originated from bacterial enrichments with and without physical supports, reflecting differences in the bacterial community composition (Fig. 2a). The type of physical support

used in the bacterial enrichments also resulted in differences, in particular dominant populations reflected by the DGGE bands. These differences were observed in terms of number and intensity of bands. Most bands observed in the profiles corresponding to the enrichment without any physical support were not observed in the enrichments with support, indicating that the bacteria related to those bands were not favored under the conditions settled for the other enrichments (Fig. 2a, blue arrows in lanes 1–3). On the other hand, profiles derived from the enrichments containing supports for an increase of biomass revealed specific bands not observed in the enrichment without any physical matrix (Fig. 2a, red and green arrows in lanes 4–12). Enrichments with arenite and shale as matrices were more similar to each other when compared to the enrichment with polyurethane foams (Fig. 2b). In addition, band patterns corresponding to the arenite and shale enrichments showed the lowest relative richness (number of bands); whereas, the enrichments with polyurethane foam showed the highest complex band patterns (Fig. 2a, lanes 4–6).

16S rRNA gene libraries, ARDRA and phylogenetic analysis

The bacterial diversity of the four different enrichment cultures evaluated in this work was determined by analysis of 16S rRNA gene clone libraries.

A total of 95 clones from the WS (without support) enrichment, 46 clones from the PF (polyurethane foams) enrichment, 63 clones from the S (shale) enrichment and 50 clones from the A (arenite) enrichment were screened by ARDRA aiming to select different ribotypes for subsequent sequencing and phylogenetic analysis. Combined data from ARDRA and sequencing analyses allowed to unravel the bacterial diversity recovered in the petroleum enrichments (Fig. 3). Clones were related to sequences available at the Genbank and RDP (Ribosomal Database Project) public database.

The ARDRA analysis of the 95 clones from the WS enrichment showed seven distinct restriction profiles. Clones were affiliated to the genera *Petrotoga* (48.4%) (Phylum Thermotogae) and *Bacillus* (51.6%) (Phylum Firmicutes) (Fig. 3a).

Fifteen distinct ribotypes were detected in the PF enrichment. Sequencing of clones representing such ribotypes revealed a more diversified microbiota, which included the genera *Rahnella* (13%), *Pseudomonas* (15.2%), *Achromobacter* (6.5%) and *Acinetobacter* (2.2%) (Phylum Proteobacteria), *Geobacillus* (13%), *Paenibacillus* (10.8%), *Thermicanus* (6.5%), *Weissella* (4.4%), *Leuconostoc* (6.5%) and *Bacillus* (2.2%) (Phylum Firmicutes), *Petrotoga* (4.4%) (Phylum Thermotogae) and *Kocuria* (8.7%) (Phylum Actinobacteria). Some clones (6.5%) were related to sequences from uncultured bacteria, and thus considered unaffiliated (Figure 3b). The ARDRA screening of the 63 clones from the S enrichment yielded three distinct ribotypes. Analyses of the clone sequences revealed that the bacterial community was composed basically by the phyla Thermotogae, represented by the genus *Petrotoga* (94%), and Proteobacteria, represented by the genus *Rahnella* (6%) (Fig. 3c).

Finally, ARDRA screening also revealed three distinct ribotypes from 50 clones recovered from the A enrichment.

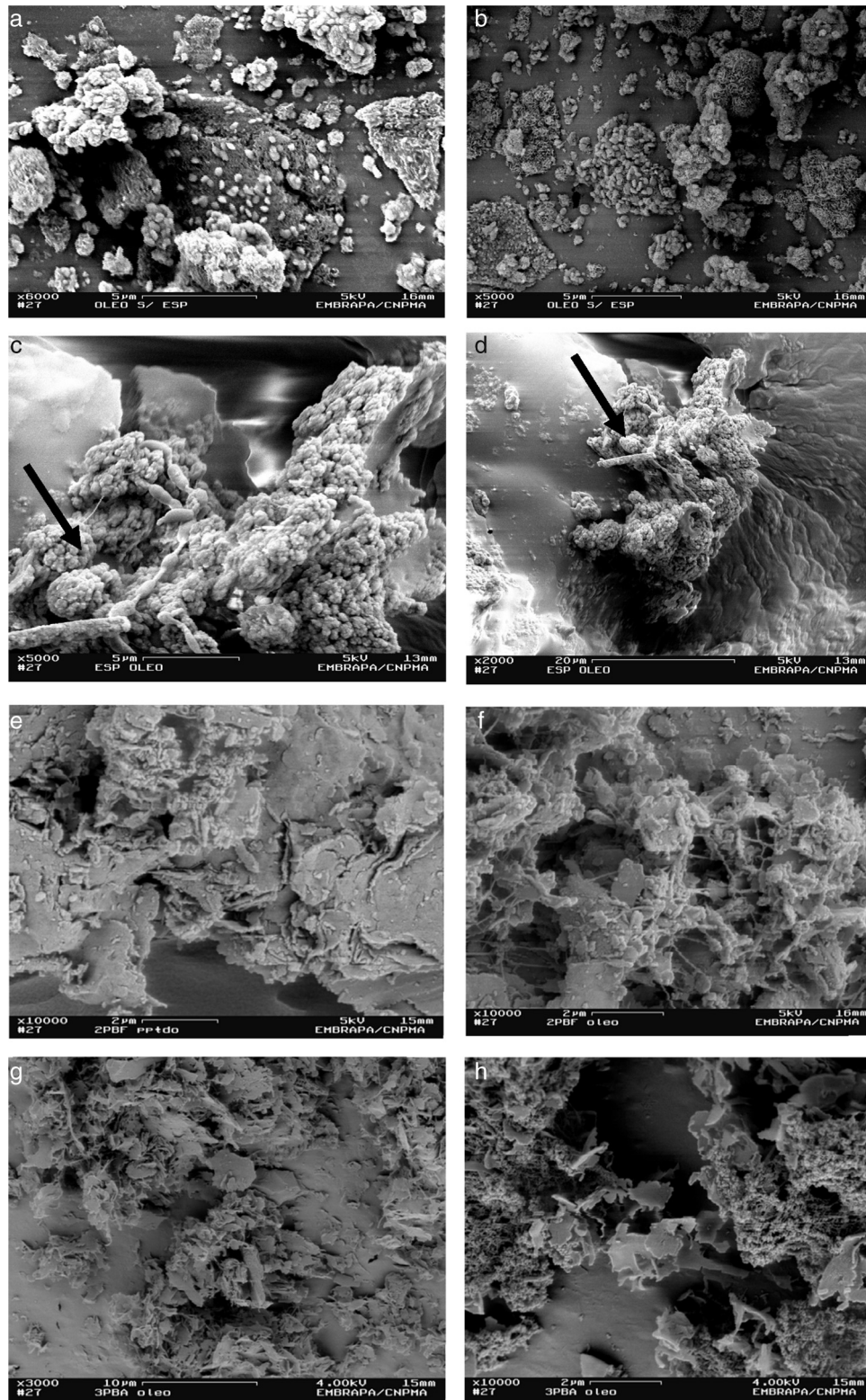


Fig. 1 – SEM analyses of different bacterial enrichments from Campos Basin oil. (a, b) Oil without support; (c, d) oil added with polyurethane foams; (e, f) oil added with shale; (g, h) oil added with arenite. Arrows indicates biofilm formation.

Similarly to the S enrichment, the experiments using arenite as physical support revealed the massive predominance of the genus *Petrotoga* (Phylum Thermotogae) (96%). Besides Thermotogae, the Phylum Firmicutes was also identified in

this microbiota, represented by clones related to the genus *Thermicanus* (4%) (Fig. 3d).

Phylogenetic analysis allowed the identification of many bacterial members at the specie level (Fig. 4). The majority

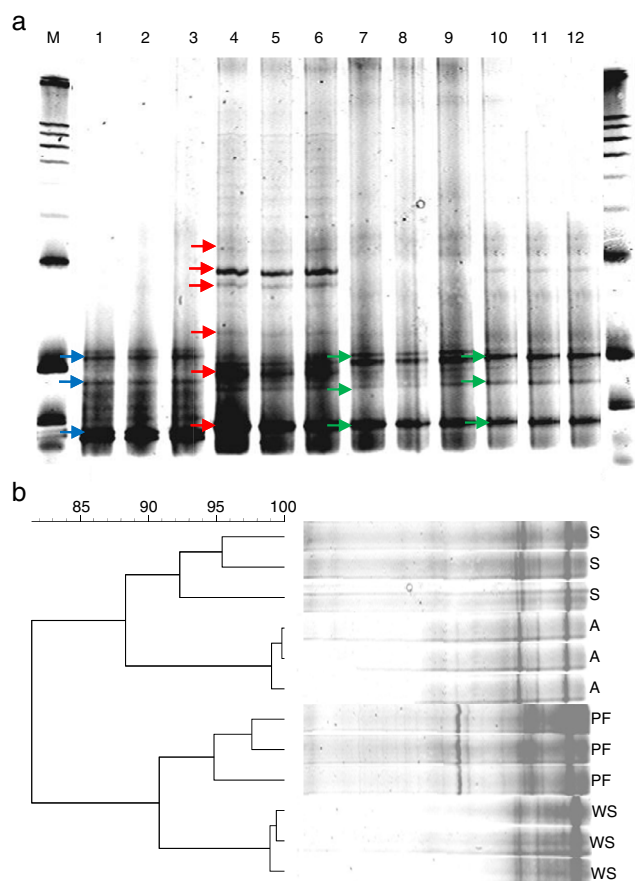


Fig. 2 – (a) DGGE analyses showing the distinct band profiles. M: marker; Lanes 1–3: WS enrichments; lanes 4–6: PF enrichments; lanes 7–9: S enrichments; lanes 10–12: A enrichments. (b) Dendrogram showing the grouping of different profiles among the enrichments and distinct matrices.

of the clones related to *Petrotoga* present in the libraries from the A, S, PF and WS enrichments were related to three *Petrotoga* species, being *Petrotoga halophila* the closest one. These clone sequences exhibited high sequence similarity (99%) with *Petrotoga* strains isolated from oil reservoirs (Table 2). Actually, the majority of the analyzed clones in all libraries were affiliated with the genus *Petrotoga*, amounting to 155 clones, 48 from A, 59 from S, 46 from WS and 2 from PF enrichment. Clones related to *Bacillus* present in the libraries from the WS (49 clones) and PF (1 clone) enrichments were related to *Bacillus ginsengihumi*, showing 99% sequence similarity (Table 2). Ten clones present in the S and PF libraries were related to *Rahnella aquatilis* and *Rahnella* sp. (sequence similarity between 80% and 99%). Five clones were closely related to the species *Thermicanus aegyptius*, two from the A library and three from the PF library, showing 99% of sequence similarity.

Other bacterial species were identified only from the PF library (Fig. 4). One clone related to *Acinetobacter ursingii* was found, showing 99% sequence similarity. Three clones were related to *Leuconostoc mesenteroides* and two clones to *Weissella confusa*, both species isolated from fermented food (Table 2),

with 99% sequence similarity. Six clones were clustered with high bootstrap value with the type strains of *Geobacillus thermoglucosidasius* and *Geobacillus caldoxylosidasius*, showing 99% sequence similarity. In this case, it was not possible to define the identification of the clones at the species level, since they were recovered in a tight cluster with both species. Five clones were closely related to *Paenibacillus naphthalenovans* (99% sequence similarity). In addition, four clones were clustered with the actinobacterium *Kocuria kristinae* (100% bootstrap value), three clones with *Achromobacter xylosoxidans* (99% sequence similarity; 100% bootstrap value) and, finally, seven clones with the type strain of *Pseudomonas putida*.

Discussion

In this study, three different types of matrices (polyurethane foam, shale and arenite) were evaluated as supports for biomass immobilization and increase in anaerobic enrichments from a biodegraded petroleum sample (Campos Basin). Shale, in particular, constitutes nearly 70% of the rocks present in a sedimentary basin. Geochemical analyses showed that almost all hydrocarbons of the petroleum from Campos Basin are from shale, belonging to the Lower Cretaceous Lagoa Feia Formation.³⁶ In this sense, this material is probably of great relevance in providing a substrate for the microbial growth in petroleum reservoirs. The use of shale in bacterial enrichment from the petroleum sample allowed the cells to grow around the shale slices generating a type of biofilm. In fact, cell aggregates, as well as the EPS structure responsible for the maintenance of cell cohesion, were observed for the enrichments employing the two other physical supports, arenite and polyurethane foams, but not for the WS (without support) enrichment. These results confirm the usefulness of these types of physical supports to enable biofilm formation and increase the microbial biomass from low cellular abundance samples, corroborating previous findings.^{28,37} However, in terms of DNA recovery, polyurethane foams were the most efficient material.

It is known in the literature that bacteria can survive in associations, named as biofilms, producing dense biomass and polymeric substances able to keep them together as a unit.³⁸ In petroleum reservoirs the presence of microbial biofilms are mostly associated with corrosion process.³⁹ In case of specific groups as sulfate reducing bacteria, for instance, it is generally accepted that souring microbiota can form mixed community biofilms on the reservoir mineral matrix.⁸

Immobilization often simulates what occurs naturally when cells grow on surfaces or within natural structures. Numerous biotechnological processes are incremented by the use of microbial immobilization techniques. These techniques can be divided into four types, based on the physical characteristics of the supports employed: (1) attachment or adsorption on solid carrier surfaces, (2) entrapment within a porous matrix, (3) self-aggregation by flocculation (natural or induced) and (4) cell containment barriers.⁴⁰ However, the choice of a material as the ideal physical support can be determinant in the selection of a microbial community,²⁸ which could be observed also in this work. The authors demonstrated that

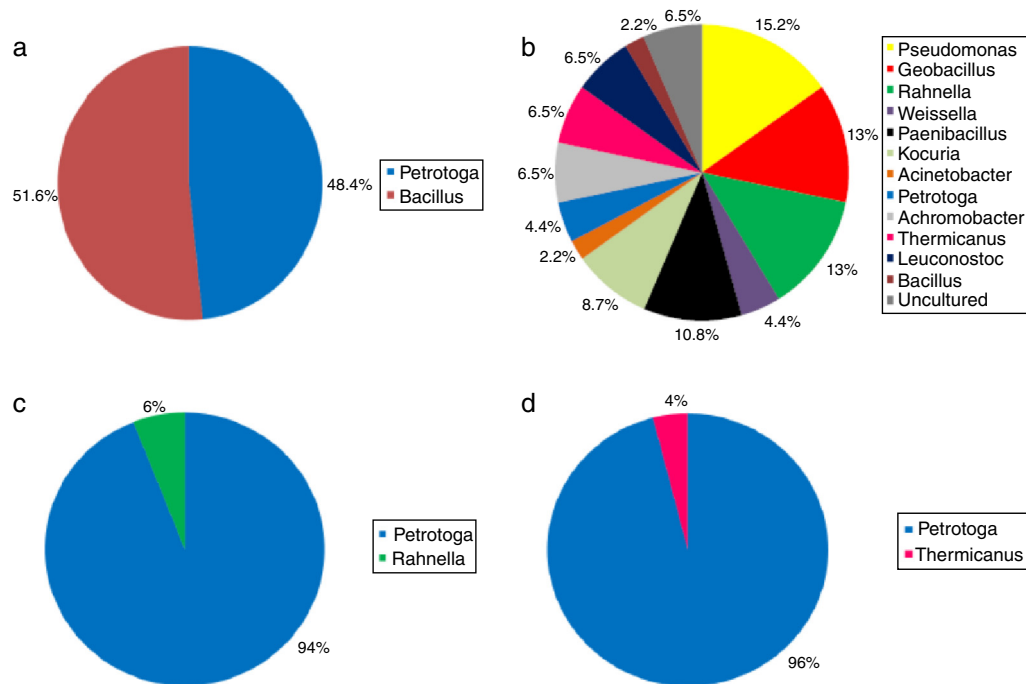


Fig. 3 – Occurrence of bacterial genera in the petroleum enrichments: (a) Enrichment without support material (WS) (95 clones, 7 ribotypes); (b) enrichment with polyurethane foams (PF) as support material (46 clones, 15 ribotypes); (c) enrichment with shale (S) as support material (63 clones, 3 ribotypes); and (d) enrichment with arenite (A) as support material (50 clones, 3 ribotypes).

Table 2 – Bacterial diversity of anaerobic enrichments from a biodegraded petroleum sample from Campos basin revealed by culture-independent methods.

Bacterial genus	Library	No. of total clones	Closest relatives	Source	% similarity
Petrotoga	A	48	<i>Petrotoga halophila</i>	Oil well	99
	S	59	<i>Petrotoga halophila</i>	Oil well	99
	PF	2	<i>Petrotoga halophila</i>	Oil well	98
	WS	46	<i>Petrotoga halophila</i>	Oil well	99
Bacillus	PF	1	<i>Bacillus ginsengihumi</i>	Spacecraft clean rooms	99
	WS	49	<i>Bacillus ginsengihumi</i>	Orchards in China/Spacecraft clean rooms	99
Thermicanus	A	2	<i>Thermicanus aegyptius</i>	Oxic soil	99
	PF	3	<i>Thermicanus aegyptius</i>	Oxic soil	99
Rahnella	S	4	<i>Rahnella aquatilis</i>	Fruits and vegetables	99
	PF	6	<i>Rahnella</i> sp.		99
Pseudomonas	PF	7	<i>Pseudomonas putida</i>	River water polluted with phenolic compounds/soil	99
Geobacillus	PF	5	<i>G. caldxylosidasius</i>	Cool soil environments	99
		1	<i>G. thermoglucosidasius</i>		99
Paenibacillus	PF	5	<i>P. naphthalenovorans</i>	Rhizosphere of salt marsh plants	99
Kocuria	PF	3	<i>K. kristinae</i>	Princess Elisabeth Station Antarctica	99
Achromobacter	PF	3	<i>A. xylooxidans</i>	Non-rhizobial plant	99
Leuconostoc	PF	1	<i>L. mesenteroides</i>	Fermented food	100
		2	<i>Leuconostoc lactis</i>	Dairy products	99
Acinetobacter	PF	1	<i>Acinetobacter</i> sp.	Spacecraft associated clean rooms	99
Weissella	PF	2	<i>Weissella confusa</i>	Fermented food	99

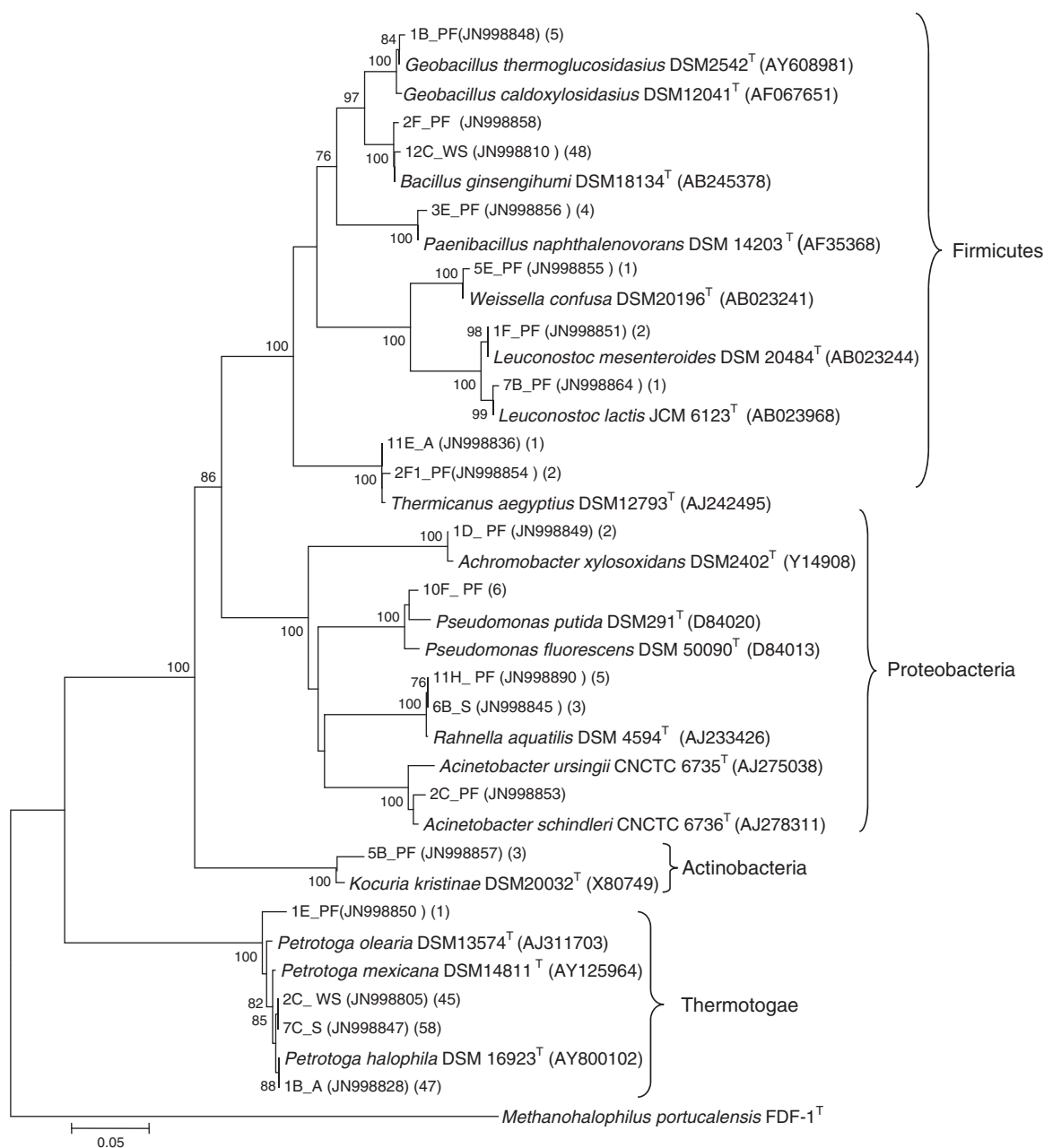


Fig. 4 – Phylogenetic analysis based on partial bacterial 16S rRNA sequences of clones from diverse enrichment samples and related species. Bootstrap values greater than 70% are listed. GenBank accession numbers are listed after species names. Numbers in brackets correspond to additional clones presenting. 97% sequence similarity with the clones represented in the tree. Letters PF, WS, A and S correspond to the sample libraries. *Methanohalophilus portucalensis* was used as outgroup.

different microbial genera were recovered depending on the type of support material used, suggesting that the appropriateness of the support material in a microbial enrichment will depend on the microbial group of interest.

The DGGE analyses were performed as an efficient tool for the comparison of the microbial diversity recovered from the different bacterial enrichments implemented in this study. The results demonstrated that the polyurethane foams allowed the development of the most complex bacterial community, reflected by the higher number of dominant populations in the DGGE profiles, differently from the

other enrichments, which yielded the lowest number of bands.

The composition of phytotypes at the phylum level was similar between the libraries derived from the WS (without support) and A (arenite) bacterial enrichments, both showing Thermotogae and Firmicutes as the predominant phyla. The phylum Thermotogae was the most predominant in the arenite library (96%), while in the WS clone library the Thermotogae and Firmicutes were found at the similar proportion (48.4% and 51.6% respectively). In many microbial community studies from oil reservoirs, injection water or other

petroleum-associated environments, members of these two phyla are common inhabitants and, not rarely, predominant.^{2,41,42} Members of the family Thermotogaceae (order Thermotogales) belong to two physiological groups: extreme thermophiles that grow at temperatures above 70 °C and moderate thermophiles that grow at lower temperatures.⁴³ The genus *Petrotoga*, as the genera *Geotoga* and *Thermotoga*, is described as occurring exclusively in petroleum reservoirs.^{44,45}

The S (shale) bacterial enrichment also yielded a community composed mainly by the phylum Thermotogae (94%), represented by the genus *Petrotoga*, followed by the phylum Proteobacteria (6%), represented only by the genus *Rahnella*. Although not commonly described in petroleum environments, this bacterium was previously reported as being involved in hydrocarbon biodegradation in Antarctic soils contaminated with polycyclic aromatic hydrocarbons (PAHs) and in enrichments from Brazilian petroleum samples.^{46,47} In a recent study of our research group,¹⁶ the bacterial diversity of the aerobic (AER) and anaerobic (ANA) enrichments of oil samples from Potiguar Basin (RN, Brazil) was evaluated by 16S rRNA clone library analysis, and 38.4% of the clones from the anaerobic enrichment were affiliated to the genus *Rahnella*.

The clone library originated from the PF (polyurethane foams) enrichments was more diverse at both genus and phylum levels. The use of polyurethane foam as support for the immobilization process has been already reported in literature, with wide application in studies of molecules such as enzymes,⁴⁸ dyes⁴⁹ and even building material.⁵⁰ Polyurethane foam presents some features such as porosity, which does not only increase the surface area but also minimize the diffusion limitation for substrate and product.⁴⁸ Silva et al.²⁸ also described the use of polyurethane foams as an efficient support material for anaerobic biomass immobilization and increment of microbial growth, especially for sulfate reducing bacteria.

Phylogeny analysis of the clone library derived from the PF enrichments revealed that the clones were affiliated to four different phyla in distinct abundances (Proteobacteria, Thermotogae, Firmicutes and Actinobacteria). Differences were more pronounced when comparing the microbiota recovered from the enrichments at the genus level. The PF enrichments allowed the recovery of representatives of 12 different genera, corroborating the efficiency of polyurethane foams for the improvement of the microbial diversity recovery. Many of the genera found using polyurethane foam as physical support, such as *Geobacillus*, *Bacillus*, *Achromobacter*, *Acinetobacter*, *Pseudomonas*, *Kocuria* and *Paenibacillus*, are described as organisms living in petroleum-associated environments, and some of them are involved in hydrocarbon degradation processes. The *Geobacillus* spp. constitute a thermophilic group, classified into the order Bacillales, described as being isolated from petroleum reservoirs.⁵¹ Liu and co-workers⁵² studied the *alkB* genes in species of this group, suggesting their potential as hydrocarbon degraders. The *Achromobacter* species have been previously described in the literature as hydrocarbon degraders and/or associated with oil field environments.^{22,53,54} A microbial diversity study conducted with injection water samples in platforms of the Campos Basin (Brazil) reported that 24% of the total 16S rRNA clones were related to the

Achromobacter genus.⁵⁵ The *Pseudomonas*, *Bacillus* and *Acinetobacter* spp. are commonly described as inhabitants of petroleum-associated environments, including reservoirs^{3,22,42,56-58} and their role is often linked to the hydrocarbon degradation process, including the production of biosurfactants.^{59,60} Similarly, the genera *Kocuria* and *Paenibacillus* have already been related to oil reservoirs⁶¹ and also to hydrocarbon degradation.^{62,63} The *Paenibacillus* spp. have been isolated from Iranian oil wells⁶¹ and the *Kocuria* from Chinese oil fields.⁶⁴

The *Leuconostoc* is a bacterial group commonly described living in fresh plants and plays an important role in several industrial and food fermentation processes.⁶⁵ Nonetheless, the presence of the genus *Leuconostoc* has already been reported in an oil field environment.²¹ Members of the genus *Weissella* are usually isolated from food and vegetables and have been involved with fermentative processes of food products.⁶⁶ Recently, Silva and co-workers¹⁶ have found *Kocuria*, *Bacillus*, *Weissella*, *Achromobacter*, *Acinetobacter* and *Leuconostoc* in two different petroleum samples (Potiguar Basin, RN) from Brazilian reservoirs.

The genus *Thermicanus* was first described as a group encompassing thermophilic, fermentative microaerophilic bacteria living in soils.⁶⁷ These bacteria were co-isolated together with a thermophilic acetogen, *Moorella thermoacetica*, from oxic soil obtained from Egypt, and these two species were shown to grow commensally on oligosaccharides via the interspecies transfer of H₂ and formate and lactate.⁶⁷ These data suggest that *Thermicanus* might be indirectly involved in the complex syntrophic degradation of hydrocarbons in oil-associated environments.

The results obtained using the different support materials, polyurethane foams, shale and arenite, revealed that all of them led to an increase of the microbial biomass, when compared to the enrichment without any physical support. However, it was expected that the use of shale and arenite would allow the recovery of a more diversified microbiota from oil samples, considering that they participate in the composition of the reservoir rock and are a natural support for biofilm formation in such environments. In fact, polyurethane foams yielded the highest microbial diversity. This could be explained by the porous nature of the polyurethane foams that allows an intense biofilm formation and EPS production, which in turn allows microorganisms that do not have affinity with the material surface to get attached to the first biofilm colonizers, thus increasing the microbial diversity.

Despite the diversity found in the anaerobic enrichments, members related to the sulfate reduction metabolism were not observed. This fact could be explained by the short period of incubation, 60 days, which could lead to an initial selection of facultative anaerobic heterotrophic bacteria, capable of degrading faster the nutrient source, becoming more abundant than sulfate-reducing members.

Conclusion

This work describes the use of three different physical supports – shale, arenite and polyurethane foams – for efficient biomass recovery in petroleum enrichments in sulfate

reducing conditions. Molecular techniques and SEM were used as tools to assess the microbial diversity as a function of the physical support employed in the enrichments. Results revealed *Petrotoga* as the most abundant genus in the enrichments using shale and arenite as physical support. On the other hand, the enrichment using polyurethane foams was more diverse and allowed the identification of 12 different bacterial genera. Finally, the combined data gathered in this work demonstrated the usefulness of physical supports for the enrichment of low abundance microorganisms found in particular environments, such as deep oil reservoirs, enabling subsequent microbiological, physiological, genomic or metagenomic analyses.

Conflict of interest

The authors declare that there is no conflict of interest.

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