Responses of microbial community from tropical pristine coastal soil to crude oil contamination

Daniel Morais, Victor Pylro, Ian M Clark, Penny R Hirsch, Marcos R Tótola

Brazilian offshore crude oil exploration has increased after the discovery of new reservoirs in the region known as pré-sal, in a depth of 7,000 m under the water surface. Oceanic Islands near these areas represent sensitive environments, where changes in microbial communities due oil contamination could stand for the loss of metabolic functions, with catastrophic effects to the soil services provided from these locations. This work aimed to evaluate the effect of petroleum contamination on microbial community shifts (Archaea, Bacteria and Fungi) from Trindade Island coastal soils. Microcosms were assembled and divided in two treatments, control and contaminated (weathered crude oil at the concentration of 30 g kg\(^{-1}\)), in triplicate. Soils were incubated for 38 days, with CO2 measurements every four hours. After incubation, the total DNA was extracted, purified and submitted for target sequencing of 16S rDNA, for Bacteria and Archaea domains and Fungal ITS1 region, using the Illumina MiSeq platform. Three days after contamination, the CO2 emission rate peaked at more than 20x the control and the emissions remained higher during the whole incubation period. Microbial alpha-diversity was reduced for contaminated-samples. Fungal relative abundance of contaminated samples was reduced to almost 40% of the total observed species. Taxonomy comparisons showed rise of the Actinobacteria phylum, shifts in several Proteobacteria classes and reduction of the Archaea class Nitrososphaerales. This is the first effort in acquiring knowledge concerning the effect of crude oil contamination in soils of a Brazilian oceanic island. This information is important to guide any future bioremediation strategy that can be required.
Responses of microbial community from tropical pristine coastal soil to crude oil contamination

Daniel K. Morais¹,²,³* Victor S. Pylro², Ian M. Clark³, Penny R. Hirsch³, Marcos R. Tótola¹*

¹Microbiology Department, Universidade Federal de Viçosa, Viçosa/MG, 36570-900, Brazil.
²Genomics and Computational Biology Group, René Rachou Research Center (CPqRR-FIOCRUZ), Belo Horizonte/MG, 30190-002, Brazil
³AgroEcology Department, Rothamsted Research, Harpenden/Herts, AL52JQ, United Kingdom.

*Correspondence: Daniel Morais and Marcos R. Tótola. Department of Microbiology, Universidade Federal de Viçosa, Av. P.H. Rolfs, SN, Viçosa, MG, 36570-900, Brazil. E-mail: kdanielmorais@hotmail.com; totola@ufv.br.
Phone: +55 31 3899-2903

ABSTRACT

Brazilian offshore crude oil exploration has increased after the discovery of new reservoirs in the region known as pré-sal, in a depth of 7.000 m under the water surface. Oceanic Islands near these areas represent sensitive environments, where changes in microbial communities due oil contamination could stand for the loss of metabolic functions, with catastrophic effects to the soil services provided from these locations. This work aimed to evaluate the effect of petroleum contamination on microbial community shifts (Archaea, Bacteria and Fungi) from Trindade Island coastal soils. Microcosms were assembled and divided in two treatments, control and contaminated (weathered crude oil at the concentration of 30 g kg⁻¹), in triplicate. Soils were incubated for 38 days, with CO₂ measurements every four hours. After incubation, the total DNA was extracted, purified and submitted for target sequencing of 16S rDNA, for Bacteria and Archaea domains and Fungal ITS1 region, using the Illumina MiSeq platform. Three days after contamination, the CO₂ emission rate peaked at more than 20x the control and the emissions remained higher during the whole incubation period. Microbial alpha-diversity was reduced for contaminated-samples. Fungal relative abundance of contaminated samples was reduced to almost 40% of the total observed species. Taxonomy comparisons showed rise of the Actinobacteria phylum, shifts in several Proteobacteria classes and reduction of the Archaea class Nitrososphaerales. This is the first effort in acquiring knowledge concerning the effect of crude oil contamination in soils of a Brazilian oceanic island. This information is important to guide any future bioremediation strategy that can be required.
INTRODUCTION

The offshore petroleum exploration offers risks to the whole sea life, as their hydrocarbons are toxic, mutagenic, teratogenic and carcinogenic (Hentati et al, 2013; Mckee et al, 2013). These toxic compounds tend to accumulate in the environment after spillage events, but factors as temperature, sun light, high exchange of gases and biological activity can remove the lighter portions of the crude oil in the first weeks after leakage. However, the recalcitrant portion of the oil stays in the environment for years (Huesemann et al., 2002; Trindade et al., 2005). The British Petroleum review of 2015, states that crude oil is still the dominant energy source in Brazil and that the consumption has kept rising since 2013. The recent discovery of crude oil reservoirs in the so-called pré-sal (pre-salt) reservoir is considered an excellent opportunity to supply the country’s economic and energetic demands (Lima, 2010), but possible oil spills events should be a major concern.

Trindade Island is located at the South Atlantic Ocean, 1,160 km from the city of Vitória, capital of Espírito Santo State, Brazil, being the closest oceanic island from these new Brazilian petroleum offshore exploration area. It hosts a peculiar and endangered biodiversity (Alves & Castro, 2006; Mohr et al., 2009), so the development of conservation approaches to maintain these unique ecosystems is required. It is well known that microbes are fundamental to several soil processes, including changes on physicochemical properties and degrading recalcitrant and toxic compounds (Elliot et al., 1996; Haritash & Kaushik, 2009). The expected scientific benefits from increasing knowledge on Trindade Island soil microbial diversity are extensive, including a better understanding of the roles played by these communities to empowering bioremediation actions.

The input of a mixture of hydrocarbons, as crude oil, directly influences the structure of microbial populations in soils (Hamamura et al., 2006). In contamination events, changes in soil properties, such as crude oil viscosity increasing, ageing, sorption of nutrients and toxicity cause the microbial community to shift towards profiting oil resistant populations. Some microorganisms are capable of degrading crude oil hydrocarbons through a number of aerobic and anaerobic metabolic pathways, using these compounds as sources of carbon and energy
(Zobell, 1946; Atlas, 1981; Haritash & Kaushik, 2009) comprising an appropriate target for studies focused on alleviating any possible impacts of soil contamination.

In the last 10 years, after the development of the Next Generation Sequencing (NGS) technology, microbial community studies have undergone a major boost (Caporaso et al., 2012; Loman and Pallen, 2015; Markowitz et al., 2015). Nevertheless, research related to crude oil contamination is primarily focused on the water column, without applying NGS (Huetel et al., 2014; Rodrigues et al., 2015), or only performed after an accidental contamination event (Lamendella et al., 2014; Rodriguez et al., 2015), lacking any proper control.

Crude oil hydrocarbons are expected to impact soil microbial communities through toxic effects of the oil components, enriching the environment with hydrocarbon degrading microorganisms. Here, we aimed to evaluate the microbial community shifts (Archaea, Bacteria and Fungi) from Trindade Island coastal soil under crude oil contamination, using state of the art NGS approach on a controlled microcosm experiment, in order to access the whole soil microbiota, including the nonculturable and low abundance ones.

MATERIAL AND METHODS

Sampling site and soil analysis
Trindade Island soil was randomly sampled, 10 soil cores with 6 cm of diameter to the depth of 0-10 cm, from the northeast shoreline of Trindade (coordinates: 20°30' S and 29° 19' W), under influence of native vegetation (*Cyperus atlanticus*). Soil cores were bulked, sieved (< 2mm) and stored at 4 °C, for 20 days, until microcosm assembly (Figure 1). The sampling expedition took place through April 2013. A total of 10 chemical variables (pH, P-rem, P, K, S, Ca²⁺, Mg²⁺, OM, N and C), plus soil texture, were assessed in the soil analysis. The protocol references and results are shown in Table 1, in the results section.

Soil treatment with crude oil
Firstly, to simulate the ageing of crude oil exposed to environmental conditions during spillage events, we heated 500 mL of crude oil to 90 °C and incubated for two hours in a fume hood. The resulting aged crude oil was a material highly viscous and difficult to work with. To obtain homogenous mixing of oil with soil, we dissolved the aged crude oil in hexane and
applied to a subsample of each experimental soil (Figure 1). Studies regarding the degradation or
e EXTRACTION OF HYDROCARBONS FROM SOIL SYSTEMS ROUTINELY USE ORGANIC SOLVENTS FOR SPIKING OF SOIL
with these hydrocarbons, and it is well known that organic solvents are harmful for native
microbial community of soil (Maliszewska-Kordybach, 1993; Brinch et al., 2002). Therefore,
hexane was also added to soils without crude oil to create a hexane-only contaminated control
stock. These hexane (and crude oil + hexane) exposed soil stocks were kept in a fume hood until
all hexane had evaporated. We then added 10 g of the control stock soil (hexane evaporated) to
the flasks corresponding to ‘Control’, and made up to 20 g with the corresponding soil that had
not been exposed to hexane. The same procedure was repeated for the stock soils contaminated
with crude oil, corresponding to the treatment ‘Crude Oil’. This combination method was
required to repopulate the native soil microbial community injured by hexane. The final
concentration of crude oil was 30 g kg^{-1}. The flasks were incubated at 26° C and the soil
moisture was kept at 60% of its water holding capacity (remoistened periodically with deionised
water upon reaching c. 50% water holding capacity).

**Experimental design**

To evaluate the effects of crude oil on the soil microbial community, we assembled six
soil microcosms in borosilicate Wheaton® (USA) respirometer flasks of 250 mL, containing 10 g
of soil (dry weight). The microcosms were incubated at 26° C for 23 days, and the microbial
activity was monitored by quantifying CO_{2} emissions every 4 hours, using a continuous-flow
respirometer coupled to an infrared CO_{2} detector (TR-RM8 Respirometer Multiplex – Sable
System) (Heinemeyer et al., 1989). After this 23 days, acclimatizing period, three flasks (3
replicates) received further 10 g of stock soil treated with hexane for Control and three flasks (3
replicates) received further 10 g of stock soil treated with crude oil and hexane, to the treatment
Crude Oil. The final concentration of the Crude Oil treatment was 30 g kg^{-1}. After the settlement
of the treatment replicates, the incubation continued for 15 more days (Figure 1). After this
period the samples were frozen using liquid nitrogen and stored at -80 °C until total community
DNA extraction.

**Molecular analyses**

*DNA extraction and quality check*
Genomic DNA was extracted and purified from each soil sample (0.5 g) using the PowerMax® Soil DNA Isolation Kit (MoBio Laboratories, Carlsbad, CA) following manufacturer’s instructions. Purity of the extracted DNA was checked using a Nanodrop ND-1000 spectrophotometer (Nanodrop Technologies, Wilmington, DE, USA) (260/280 nm ratio) and DNA concentration was determined using Qubit® 2.0 fluorometer and dsDNA BR Assay kit (Invitrogen™). Integrity of the DNA was confirmed by electrophoresis in a 0.8 % agarose gel with 1 X TAE buffer.

**High-throughput sequencing**

Sequencing was done on the Illumina MiSeq® platform (Caporaso et al. 2012) at the High-throughput Genome Analysis Core (HGAC), Argonne National Laboratory (Illinoi, USA). Bacterial and archaeal 16S rRNA genes were amplified using primers 515F (5’-GTGCCAGCMGCCGCGGTAA-3’) and 806R (5’-GGACTACHVGGGTWTCTAAT-3’) for paired-end microbial community (Caporaso et al., 2011). Fungal ITS1 region was amplified using primers ITS1F (5’-CTTGGCCATTAGAGGAAGTAA-3’) and ITS2 (5’-GCTGCGTTCTTCATCGATGC-3’) using the method described by Smith & Peay (2014).

**Data analysis**

We applied the 16S and ITS bioinformatics pipeline recommended by the Brazilian Microbiome Project, available at http://brmicrobiome.org (Pylro et al, 2014). Briefly, this pipeline uses QIIME (Caporaso et al., 2010) and Usearch 7.0 (Edgar, 2010) for filtering low quality sequences, clustering sequences of high similarity, diversity analysis, diversity comparisons and graphical plotting. For fungal ITS analysis we also used the software ITSx (Bengtsson-Palme, 2013) for taxonomic assignment improvement. The sequencing depth can affect alpha and beta diversity analysis, therefore, we used the strategy of rarefaction (randomly sub-sampling of sequences from each sample) to equalize the number of sequences per sample and to evaluate the sufficiency of the sequencing effort. We also used the Good’s coverage (Good, 1953) index to assess the coverage reached using the rarefaction level chosen. The microbial diversity changes were measured using the alpha diversity metrics: PD whole tree (for 16S rRNA gene only), Chao (Chao, 1984) and observed species. For beta-diversity estimations, we generated distance matrixes using the phylogenetic method weighted unifrac (Lozupone et al,
2005) for 16S rRNA gene sequences and the Bray-Curtis (Bray & Curtis, 1957) method for ITS sequences. We plotted the beta-diversity distance matrixes using a bi-dimensional Principal Coordinates Analysis (PCoA) and the clusters were evaluated using the cluster quality analysis (cluster_quality.py script on QIIME) (Caporaso et al., 2010), calculating the ratio of mean “distances between samples from different clusters” to mean “distances between samples from the same cluster”. The hypothesis testing method used to compare taxonomic differences between treatments was made using the bioconductor EdgeR package (Robinson, et al., 2010). The count matrix was normalized through the relative log expression (RLE) proposed by Anders and Huber (2010), where the median count is calculated from the geometric mean of all columns and the median ratio of each sample to the median library is used as the scale factor. The p-values were corrected using the Benjamining-Hochberg false discovery rate method (FDR). The R script used in this analysis is described and available at http://github.com/kdanielmorais.

RESULTS

Soil respiration and physicochemical characteristics

The Trindade Island soil physicochemical properties are listed at Table 1. The first 23 days of incubation didn’t show any difference in CO₂ emissions between the 6 microcosms. Differences were detected only at the 24º day, after the definition of the treatments (Figure 1). CO₂ emission rate of oil-contaminated samples increased 8x compared to the control in the first 4 hours (Figure 2). Three days after contamination, emission rate peaked at more than 20 times the control. CO₂ emission of the oil treated samples was higher than the control from the definition of the treatments (24th day of incubation) until the sampling of the DNA (38th day). At the last day of incubation (38th), CO₂ emission rate of the contaminated treatment was still almost 4 times higher than the control (figure 1).

Sequencing output

A total of 314,748 joined and quality filtered 16S rRNA gene Illumina® barcoded reads, and 424,269 single end quality filtered fungal ITS Illumina® barcoded reads were obtained from the soil samples (Table S1). The oil-contaminated treatment yielded a smaller number of sequences. To minimize the effects of sequencing depth variation on diversity analysis and taxa
comparison, we applied the rarefaction method (random subsampling of sequences). Estimates of alpha and beta-diversity were based on evenly rarefied OTU matrices (45,695 sequences per sample for Bacteria and Archaea and 25,315 sequences per sample for Fungi).

Diversity comparisons

The alpha diversity indexes used in this experiment represent species richness (Tables 2 and 3). We compared treatment’s effects over Bacteria/Archaea community using the estimators Faith’s PD (phylogenetic measure of diversity based on total branch length of phylogeny captured by a sample, proposed by Faith, 1992), the Chao-1 (estimator of total species richness proposed by Chao, 1984), and observed species (number of species detected) (table 2). The effects on Fungal community was measured using only the Chao-1 and Observed species estimators, as there was not an ITS1 phylogenetic tree available to use the Faith’s PD estimator. All metrics yielded similar results for Bacteria/Archaea and Fungi. The comparison between the two treatments shows a significant reduction of diversity upon the addition of oil for Bacteria, Archaea and Fungi. The fungal community was the most sensitive group to the oil addition, showing a reduction of ~40% for the indexes Chao1 and Observed species (table 3).

The rarefaction analysis (Figures 3 A and B), which plots the operational taxonomic unit (OTU) richness as a function of sequencing depth, and the Good’s coverage shows that sequencing effort was sufficient to capture the Bacterial, Archaeal and Fungal diversity of samples. The analysis also confirms that crude oil had a reductive effect on microbial diversity.

The beta diversity analysis was performed using (Figure 4) Weighted Unifrac for 16s rRNA gene and Bray-Curtis for fungal intergenic spacer ITS1 due to the lack of a phylogenetic tree for ITS1 marker. Both methods showed two very distinct clusters separating the treatments Control and Crude Oil (Cluster quality. 16S = 2.36 and ITS = 2.14).

Taxonomic comparison

The taxonomic distributions of Bacteria/Archaea are shown in Figure 5 at phylum level. The control treatment show 6% of sequences to be from the Archaea domain, 93.4% from Bacteria domain and 0.5% were not asigned to any taxa from the GreenGenes database (DeSantis et al., 2006). For Archaea, we found only three representatives: the species Candidatus nitrosophaere belonging to the phylum Crenarchaeota, the order E2 belonging to the phylum
Euryarchaeota and the order YLA114, belonging to the phylum Parvarchaeota. The addition of oil reduced the relative abundance of Archaea to 2.7%.

We identified 225 orders in the bacterial group of the control samples. The most abundant bacterial orders in the control were Acidobacteria order iii1-15 (7%), Rhizobiales (6.5%), Rubrobacterales (6.3%), Nitrospherales (6.1%), Xanthomonadales (4.8%), Syntrophobacterales (4.2%), Gaielalles (4%) and Myxococcales (4%). Oil-contaminated samples presented 224 orders, and the most abundant orders were Actinomycetales (17%), Acidobacteria order iii1-15 (8.5%), Rhizobiales (6.4%), Burkholderiales (4%), Xanthomonadales (3.9%), Chloroacidobacteria order RB41 (3.4%), Sphingomonadales (3%), Acidimicrobiales (2.9%). The abundance of 57 taxa was significantly different between Control and Crude Oil (Table 4).

Fungal taxonomy analysis (Figure 6) was assessed using the UNITE database version 7 (Kõljalg et al., 2005). 5% of the reads from non-contaminated soil were not assigned to any taxonomic group. For the crude oil treatment, only 0.7% of the sequences did not match to a taxon. We found 29 orders in the fungal group of the control samples. The most abundant orders in the control were Hypocreales (41%), Mortierellales (27%) and Sordariales (7.5%). Oil-contaminated samples presented 29 orders, and the most abundant orders were Mortierellales (70%), Hypocreales (24%) and Botryosphaeriales (1.1%). Abundance of 6 taxa was significantly different between control and oil contaminated soils (Table 5).

**Discussion**

In this study, we have applied high throughput sequencing to evaluate the effect of crude oil contamination on Trindade Island soil microbiota. We found that crude oil had a deleterious effect on microbial alpha-diversity (Tables 2 and 3). This result is similar to the obtained by Yang (2014), as crude oil was thought to have an eco-toxicological effect. The higher amount of CO₂ evolved in the crude oil treated-soil (Figure 2) is related to the oil stressing effect (Franco et al., 2004), and the further peaks observed in the Figure 2, might be related to different fractions of oil being degraded according to its bioavailability.

Despite the toxic effect, some taxa are able to use oil hydrocarbons as a source of carbon and energy being favoured by oil amendment, and gradually overcoming the populations lacking those abilities. The effects on fungal diversity were more marked than that observed on prokaryotic diversity, corroborating Embar (2006), who reported a rapid increase in abundance
and shift in diversity in the fungal community in response to oil contamination. The strong effect of oil on the fungal diversity may also be explained by metabolic differences between eukaryotes and prokaryotes. This effect relates to the increased toxicity of polycyclic aromatic hydrocarbons, present in crude oil, after metabolic activation mediated by the enzyme cytochrome P450 (CYP) of eukaryotes. The majority of carcinogens in the environment are inert by themselves and require the metabolic activation by CYP, in order to exhibit carcinogenicity (Shimada and Fuji-Kuriama, 2003). The CYP genes belong to the superfamily of dioxygenases, present in all domains of life. Genes that code for dioxygenases in prokaryotes are related to toxin and xenobiotic degradation, while in eukaryotes CYP genes may be related to a plethora of functions, ranging from biosynthesis of hormones to chemical defence in plants (Werck-Reichhard et al., 2000).

We observed the formation of two distinct clusters representing the control samples and the crude oil contaminated samples during the analysis of beta-diversity (Figure 4). We found that bacterial/archaeal oil-contaminated replicates showed a broader spread in the PCoA, while oil-contaminated replicates in fungal communities are clustered more tightly (figure 4B). Because of selective pressure, the taxa resistant to the contamination event and the populations able to degrade hydrocarbons will gradually outnumber the rest of the community in the curse of succession (Yang et al., 2014). Therefore, as oil presented a toxic effect, we would expect that the bacterial community of contaminated samples would show a more compact clustering, as happened with the fungal community. However, as the bacterial community comprises c. 30x more OTU than the fungal, the shifts in the bacterial relative abundance might be more related to soil microhabitats present in each replicate, than with the oil toxic effects. This phenomenon was previously observed (Juck et al., 2000; Liang et al., 2011; Yang et al., 2014), and could be explained by the appearance of new niches in the contaminated soil with further fulfilment of these niches by previously not detected (low abundance) taxa.

Soil is the most diverse environment on earth (Vogel et al., 2009), and many of the native microorganisms possess the ability to resist and degrade crude oil hydrocarbons (Franco et al., 2004; Head et al., 2006). In this study, we detected the relative abundance community shifts in Actinobacteria, Proteobacteria, Firmicutes and Planctomycetes. The phylum Actinobacteria had its abundance increased in response to crude oil addition. We detected shifts in one unidentified species from the family Streptomycetaceae, one specie from the genus Streptomyces and one
species from the order Solirubrobacterales. Interestingly, the genera Nocardia represented less than 0.01% of the total sequences in the control samples and shifted to 9.4% of the sequences in the crude oil samples (Table 4). Several studies have reported Actinobacteria as a good option for removing recalcitrant hydrocarbon, since they are known for the production of extracellular enzymes that degrade a wide range of complex hydrocarbons. Also, many species of Actinobacteria are able to produce biosurfactants that enhance hydrocarbons solubility and bioavailability (Pizzul et al., 2007; Kim & Crowley, 2007; Balachandram et al., 2012; da Silva et al., 2015). The Actinobacteria phylum is recognized as the main alkane degrader in polar soils (Aislabie et al., 2006), besides producing multiple types of antifungals, antivirals, antibiotics, immunosuppressives, anti-hypertensives and antitumorals (Benedict, 1953; Omura et al., 2001; Khan et al., 2011; de Lima Procópio et al., 2012). Rodriguez et al. (2015) reported a significant rise in Gamma and Alphaproteobacteria relative abundance from beach sand of Florida coast, in response to the crude oil plume from the Deepwater Horizon Drilling rig accident in the Gulf of Mexico. Although some works have reported prevalence of Gram-negative bacteria upon soils contaminated with heavily weathered petroleum (Kaplan & Kitts, 2004), our work shows a big shift on Gram-positive Actinobacteria. Our results also corroborates with Chikere et al. (2009) who reports the prevalence of Actinobacteria after oil addition using cultivation dependent techniques. Grosshard et al. (2004) detected the inhibition of several proteobacteria by actinomycete strains isolated from the German Wadden sea, while Burgess et al. (1999) report that, antibiotic production may be triggered by several factors as presence of chemical substances, substrate availability, population density and many others.

We did not detect a shift in the general relative abundance of the Proteobacteria phylum (Figure 4) but, the relative abundance of the classes inside this phylum showed a significant change (Table 5). Alpha and Deltaproteobacteria classes had a major relative abundance reduction in the contaminated samples. The reduction of these two classes might even be connected, considering that, the Alphaproteobacteria with the biggest reduction was a member of the Rhodospirillales order, which is composed mainly by purple non sulphur photosynthetic microorganisms. This group fix carbon using hydrogen as an electron donor, and the member of the Deltaproteobacteria phylum that suffered the biggest reduction belongs to the Syntrophobacteraceae family, a family known for releasing H₂ as a product of organic acids fermentation. This ecological interaction is called syntrophy (McInerney et al., 1981), and its
presence could be happening as both groups were reduced by c. 50%. The Beta and Gammaproteobacteria classes rose in their relative abundance in response to crude oil treatment. The member of the Betaproteobacteria class with the biggest increase belonged to the family Comamonadaceae, this family is known by its heterotrophic denitrification capability (Khan et al., 2002) using organic compounds as electron donors. The only Archaea species we detected, *Nitrososphaera*, is an autotrophic ammonia-oxidizer (Mußmann et al., 2011) and represented 6.1% of the total sequences in the control. In the crude oil contaminated treatment this relative abundance was reduced to 2.8%. Urakawa et al. (2012) evaluating the responses of ammonia-oxidizing Archaea and Bacteria to crude oil hydrocarbons, showed that Archaea are several times more sensitive than Bacteria. The reduction of this Archaea and the increase in the relative abundance of the Comamonadaceae family individual (Table 5), mentioned above, reinforces the hypothesis raised to explain the broader cluster observed in Bacteria beta-diversity (Figure 4A).

This phenomenon was not observed for Fungi, as in the control samples we were able to detect 12 well distributed classes and in the contaminated samples, with more than 95% of the sequences belonged to the classes Sordariomycetes and Incertae. The Incertae class presented only the genera *Mortierella* and its relative abundance in the contaminated samples reached 70.3%. *Mortierella* is a Zygomycota and is known as an oleaginous microorganism, it accumulates lipids and has even been used as a strategy for biodiesel production (Ratledge, 2002; Kumar et al., 2011).

This is the first study reporting the effect of crude oil contamination in soils of the Trindade Island, a Brazilian oceanic island threatened by possible oil spills due petroleum exploration. Our results reinforces the importance of microbial diversity analysis in insulated environments, pointing out the impact of crude oil on microbial communities shifts from unexplored environments. Moreover, these finds indicate the biotechnological potential of degrading hydrocarbons soil microorganisms, fostering further studies aiming to relieve any oil contamination occurrence on Trindade Island.

**ACKNOWLEDGMENTS**

We would like to thank the Brazilian Navy and the Captain Rodrigo Otoch Chaves for the logistic support while collecting samples and Dr. Marc Redmile-Gordon (Centre for Sustainable
Soils and Grassland Systems, Rothamsted Research, UK) for the review of written English in the manuscript.

REFERENCES

Aislabie, J., Saul, D. J., Foght, J. M. (2006) Bioremediation of hydrocarbon contaminated polar soils. Extremophiles 10: 171–179.

Alves, R. J. V., & Castro, J. W. (2006) Ilhas oceânicas brasileiras: da pesquisa ao manejo. MMA Secretaria de Biodiversidade e Florestas. Brasília, DF, Brasil. 298 p.

Anders, S., Huber, W. (2010). Differential expression analysis for sequence count data Genome Biology 11, R106.

Atlas, R. M. (1981). Microbial degradation of petroleum hydrocarbons: an environmental perspective. Microbiological reviews, 45(1), 180.

Balachandran, C., Duraipandiyan, V., Balakrishna, K., & Ignacimuthu, S. (2012). Petroleum and polycyclic aromatic hydrocarbons (PAHs) degradation and naphthalene metabolism in Streptomyces sp.(ERI-CPDA-1) isolated from oil contaminated soil. Bioresource Technology, 112, 83-90.

Benedict, R. G. (1953). Antibiotics produced by actinomycetes. The Botanical Review, 19(5), 229-320.

Bengtsson-Palme, J., Ryberg, M., Hartmann, M., Branco, S., Wang, Z., Godhe, A., Nilsson, R. H. (2013). Improved software detection and extraction of ITS1 and ITS2 from ribosomal ITS sequences of fungi and other eukaryotes for analysis of environmental sequencing data. Methods in Ecology and Evolution, 4(10), 914-919.

Brinch, U. C., Ekelund, F., & Jacobsen, C. S. (2002). Method for spiking soil samples with organic compounds. Applied and Environmental Microbiology, 68(4), 1808-1816.

British Petroleum; BP statistical review of world energy. BP, London, UK. 2014. Accessible at: bp.com/statisticalreview

Burgess, J. G., Jordan, E. M., Bregu, M., Mears-Spragg, A., & Boyd, K. G. (1999). Microbial antagonism: a neglected avenue of natural products research. Journal of Biotechnology, 70(1), 27-32

Caporaso, J. G., Kuczynski, J., Stombaugh, J., Bittinger, K., Bushman, F. D., Costello, E. K., Fierer, N., Peña, A. G., Goodrich, J. K., Gordon, J. I., Huttenhower, G. A., Kelley, S. T., Knights, D., Koenig, J. E., Ley, R. E., Lozupone, C. A., McDonald, D., Muegge, B. D., Pirrung, M., Reeder, J., Sevinsky, J. R., Turnbaugh, P. J., Walters, W. A., Widmann, J., Yatsunenko, T., Zaneveld, J., Knight, R. (2010). QIIME allows analysis of high-throughput community sequencing data. Nature methods, 7(5), 335-336.
Caporaso, J. G., Lauber, C. L., Walters, W. A., Berg-Lyons, D., Huntley, J., Fierer, N., Owens, S. M., Betley, J., Fraser, L., Bauer, M., Gormley, N., Gilbert, J. A., Smith, G., Knight, R. (2012). Ultra-high-throughput microbial community analysis on the Illumina HiSeq and MiSeq platforms. The ISME journal, 6(8), 1621-1624.

Caporaso, J. G., Lauber, C. L., Walters, W. A., Berg-Lyons, D., Lozupone, C. A., Turnbaugh, P. J., Fierer, N., Knight, R. (2011). Global patterns of 16S rRNA diversity at a depth of millions of sequences per sample. Proceedings of the National Academy of Sciences, 108(Supplement 1), 4516-4522.

Chikere, C. B., Okpokwasili, G. C., & Chikere, B. O. (2009). Bacterial diversity in a tropical crude oil-polluted soil undergoing bioremediation. African Journal of Biotechnology, 8(11).

da Silva, F. S. P., Pylro, V. S., Fernandes, P. L., Barcelos, G. S., Kalks, K. H. M., Schaefer, C. E. G. R., Tótola, M. R. (2015). Unexplored Brazilian oceanic island host high salt tolerant biosurfactant-producing bacterial strains. Extremophiles, 19(3), 561-572.

de Lima Procópio, R. E., da Silva, I. R., Martins, M. K., de Azvedo, J. L., & de Araújo, J. M. (2012). Antibiotics produced by Streptomyces. The Brazilian Journal of infectious diseases, 16(5), 466-471.

DeSantis, T. Z., Hugenholtz, P., Larsen, N., Rojas, M., Brodie, E. L., Keller, K., Huber, T., Dalevi, D. Hu, F. Andersen, G. L. (2006). Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Applied and environmental microbiology, 72(7), 5069-5072.

Edgar, RC (2010) Search and clustering orders of magnitude faster than BLAST, Bioinformatics 26(19), 2460-2461.

Edwards, R. A., Rodriguez-Brito, B. Wegley, L. Haynes, M., Breitbart, M., Peterson, D. M., Saar, M. O., Alexander, S. Alexander, E. C., Rohwer, F. (2006). "Using pyrosequencing to shed light on deep mine microbial ecology". BMC Genomics 7: 57.

Elliott, L. F., Lynch, J. M., Papendick, R. I., Stotzky, G., Bollag, J. M. (1996). The microbial component of soil quality. Soil biochemistry Volume 9, 1-21.

Embar, K., Forgacs, C., & Sivan, A. (2006). The role of indigenous bacterial and fungal soil populations in the biodegradation of crude oil in a desert soil. Biodegradation, 17(4), 369-377.

Faith, D. P. (1992). Conservation evaluation and phylogenetic diversity. Biological conservation, 61(1), 1-10.

Franco, I., Contin, M., Bragato, G., & De Nobili, M. (2004). Microbiological resilience of soils contaminated with crude oil. Geoderma, 121(1), 17-30.

Grossart, H. P., Schlingloff, A., Bernhard, M., Simon, M., & Brinkhoff, T. (2004). Antagonistic activity of bacteria isolated from organic aggregates of the German Wadden Sea. FEMS microbiology ecology, 47(3), 387-396.
Hamamura, N., Olson, S. H., Ward, D. M., & Inskeep, W. P. (2006). Microbial population dynamics associated with crude-oil biodegradation in diverse soils. Applied and Environmental Microbiology, 72(9), 6316-6324.

Haritash, A. K., & Kaushik, C. P. (2009). Biodegradation aspects of polycyclic aromatic hydrocarbons (PAHs): a review. Journal of hazardous materials, 169(1), 1-15.

Head, I. M., Jones, D. M., & Röling, W. F. (2006). Marine microorganisms make a meal of oil. Nature Reviews Microbiology, 4(3), 173-182.

Heinemeyer, O., Insam, H., Kaiser, E. A., & Walenzik, G. (1989). Soil microbial biomass and respiration measurements: an automated technique based on infra-red gas analysis. Plant and soil, 116(2), 191-195.

Hentati, O., Lachhab, R., Ayadi, M., & Ksibi, M. (2013). Toxicity assessment for petroleum-contaminated soil using terrestrial invertebrates and plant bioassays. Environmental monitoring and assessment, 185(4), 2989-2998.

Hoeft, R. G., Walsh, L. M., and D. R. Keeney. 1973. Evaluation of various extractants for available soil sulfur. Soil Sci. Soc. Am. Proc. 37: 401-404.

Huettel, M., Berg, P., Kostka, J. E. (2014). Benthic exchange and biogeochemical cycling in permeable sediments. Marine Science, 6.

Juck, D., Charles, T., Whyte, L., Greer, C. (2000) Polyphasic microbial community analysis of petroleum hydrocarbon-contaminated soils from two northern Canadian communities. FEMS Microbiology Ecology 33: 241–249

Kaplan, C. W., & Kitts, C. L. (2004). Bacterial succession in a petroleum land treatment unit. Applied and Environmental Microbiology, 70(3), 1777-1786.

Khan, S. T., Horiba, Y., Yamamoto, M., & Hiraishi, A. (2002). Members of the family Comamonadaceae as primary poly(3-hydroxybutyrate-co-3-hydroxyvalerate)-degrading denitrifiers in activated sludge as revealed by a polyphasic approach. Applied and environmental microbiology, 68(7), 3206-3214.

Khan, S. T., Komaki, H., Motohashi, K., Kozone, I., Mukai, A., Takagi, M., & Shinya, K. (2011). Streptomyces associated with a marine sponge Haliclonia sp.; biosynthetic genes for secondary metabolites and products. Environmental microbiology, 13(2), 391-403.

Kim, J. S., & Crowley, D. E. (2007). Microbial diversity in natural asphalts of the Rancho La Brea Tar Pits. Applied and environmental microbiology, 73(14), 4579-4591.

Kõljalg, U., Larsson, K. H., Abarenkov, K., Nilsson, R. H., Alexander, I. J., Eberhardt, U., Erland, S., Hoiland, K., Kjoller, R., Larsson, E., Pennanen, T., Sen, R., Taylor, A. F., Tedersoo, L., Vrålstad, T., Ursing, B. M. (2005). UNITE: a database providing web based methods for the molecular identification of ectomycorrhizal fungi. New Phytologist, 166(3), 1063-1068.
Kostka, J. E., Prakash, O., Overholt, W. A., Green, S. J., Freyer, G., Canion, A., Delgardio, J., Norton, N., Hazen, T. C., Huettel, M. (2011). Hydrocarbon-degrading bacteria and the bacterial community response in Gulf of Mexico beach sands impacted by the Deepwater Horizon oil spill. Applied and environmental microbiology, 77(22), 7962-7974.

Kumar, I., Ramalakshmi, M. A., Sivakumar, U., Santhanakrishnan, P., & Zhan, X. (2011). Production of microbial oils from Mortierella sp for generation of biodiesel livestock. African Journal of Microbiology Research, 5(24), 4105-4111.

Liang, Y., Van Nostrand, J. D., Deng, Y., He, Z., Wu, L., Zhang, X., Li, G., Zhou, J. (2011). Functional gene diversity of soil microbial communities from five oil-contaminated fields in China. The ISME Journal 5: 403–413.

Lima, J. A. M. A energia que vem do mar: a herança energética do mar brasileiro. Cienc. Cult. [online], v.62, n.3, p.25-8, 2010.

Loman, N. J., & Pallen, M. J. (2015). Twenty years of bacterial genome sequencing. Nature Reviews Microbiology.

Maliszewska-Kordybach, B. (1993). The effect of temperature on the rate of disappearance of polycyclic aromatic hydrocarbons from soils. Environmental pollution, 79(1), 15-20.

Markowitz, V. M., Chen, I. M. A., Chu, K., Pati, A., Ivanova, N. N., & Kyrpides, N. C. (2015). Ten Years of Maintaining and Expanding a Microbial Genome and Metagenome Analysis System. Trends in microbiology, 23(11), 730-741.

McInerney, M. J., Bryant, M. P., Hespell, R. B., & Costerton, J. W. (1981). Syntrophomonas wolfei gen. nov. sp. nov., an anaerobic, syntrophic, fatty acid-oxidizing bacterium. Applied and Environmental Microbiology, 41(4), 1029-1039.

McKee, R. H., Schreiner, C. A., Nicolich, M. J., & Gray, T. M. (2013). Genetic toxicity of high-boiling petroleum substances. Regulatory Toxicology and Pharmacology, 67(2), S75-S85.

Mohr, L. V., Castro, J. W. A., Costa, P. M. S., & Alves, R. J. V. (2009). Ilhas oceânicas brasileiras: da pesquisa ao manejo.

Mueller, M., & Wallace, R. (2008). Enabling science and technology for marine renewable energy. Energy Policy, 36(12), 4376-4382.

Mussmann, M., Brito, I., Pitcher, A., Damsté, J. S. S., Hatzenpichler, R., Richter, A., Nielsen, J. L., Müller, A., Damis, H., Wagner, M., Head, I. M. (2011). Thaumarchaeotes abundant in refinery nitrifying sludges express amoA but are not obligate autotrophic ammonia oxidizers. Proceedings of the National Academy of Sciences, 108(40), 16771-16776.

Ōmura, S., Ikeda, H., Ishikawa, J., Hanamoto, A., Takahashi, C., Shinose, M., Takahashi, Y., Horikawa, H., Nakazawa, H., Osono, T., Kikuxhi, H., Shiba, T., Sakaki, Y, Hattori, M. (2001). Genome sequence of an industrial microorganism Streptomyces avermitilis: deducing the ability...
of producing secondary metabolites. Proceedings of the National Academy of Sciences, 98(21), 12215-12220.

Pizzul, L., del Pilar Castillo, M., & Stenström, J. (2007). Effect of rapeseed oil on the degradation of polycyclic aromatic hydrocarbons in soil by Rhodococcus wratislaviensis. International biodeterioration & biodegradation, 59(2), 111-118.

Rahman, K. S. M., Thahira-Rahman, J., Lakshmanaperumalsamy, P., & Banat, I. M. (2002). Towards efficient crude oil degradation by a mixed bacterial consortium. Bioresource technology, 85(3), 257-261.

Ratledge, C. (2002). Regulation of lipid accumulation in oleaginous microorganisms. Biochemical Society Transactions, 30(6), 1047-1049.

Robinson, M. D., McCarthy, D. J. & Smyth, J. K. (2010). edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics, 26(1): 139–140.

Rodriguez-R, L. M., Overholt, W. A., Hagan, C., Huettel, M., Kostka, J. E., & Konstantinidis, K. T. (2015). Microbial community successional patterns in beach sands impacted by the Deepwater Horizon oil spill. The ISME journal, 9(9):1928-40.

Roesch, L. F., Fulthorpe, R. R., Riva, A., Casella, G., Hadwin, A. K., Kent, A. D., Daroub, S. H., Camargo, F. A., Farmerie, W. G., Triplett, E. W. (2007). Pyrosequencing enumerates and contrasts soil microbial diversity. The ISME journal, 1(4), 283-290.

Seabra, A. A. D., Freitas, G. P. D., Polette, M., & Casillas, T. (2011). The promising oil province of the pre-salt. Revista Direito GV, 7(1), 57-74.

Shimada, T., & Fujii Kuriyama, Y. (2004). Metabolic activation of polycyclic aromatic hydrocarbons to carcinogens by cytochromes P450 1A1 and 1B1. Cancer science, 95(1), 1-6.

Smith, D. P., & Peay, K. G. (2014). Sequence depth, not PCR replication, improves ecological inference from next generation DNA sequencing. PLoS One, 9(2), e90234.

Urakawa, H., Garcia, J. C., Barreto, P. D., Molina, G. A., & Barreto, J. C. (2012). A sensitive crude oil bioassay indicates that oil spills potentially induce a change of major nitrifying prokaryotes from the Archaea to the Bacteria. Environmental pollution, 164, 42-45.

Venkateswaran, K., Iwabuchi, T., Matsui, Y., Toki, H., Hamada, E., & Tanaka, H. (1991). Distribution and biodegradation potential of oil-degrading bacteria in North Eastern Japanese coastal waters. FEMS Microbiology Letters, 86(2), 113-121.

Vogel, T. M., Hirsch, P., Simonet, P., Jansson, J. K., Tiedje, J., van Elsas, J. D., Bailey, M. J. Nalin, R., Philipot, L. (2009). TerraGenome: a consortium for the sequencing of a soil metagenome. Nature Reviews Microbiology, 7, 252.

Werck-Reichhart, D., & Feyereisen, R. (2000). Cytochromes P450: a success story. Genome Biol, 1(6), 3003-1.
Yang, S., Wen, X., Zhao, L., Shi, Y., & Jin, H. (2014). Crude oil treatment leads to shift of bacterial communities in soils from the deep active layer and upper permafrost along the China-Russia Crude Oil Pipeline route. PloS one, 9(5).

Zobell, C. E. (1946). Action of microorganisms on hydrocarbons. Bacteriological reviews, 10(1-2), 1.

**Legends**

**Tables**

Table 1 – Summary of physicochemical data for surface soil cores (0 - 10 cm) sampled at the northeast coast of Trindade Island - Brazil.

Table 2 – Average (n = 3) alpha-diversity comparison between the treatments control and crude oil for Bacteria and Archaea groups.

Table 3 – Average (n = 3) alpha-diversity comparison between the treatments control and crude oil for Fungi.

Table 4 – Bacterial/Archaeal OTUs presenting an average absolute abundance significantly different between the treatments “Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

Table 5 – Fungal OTUs presenting an average absolute abundance significantly different between the treatments “Crude Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

**Supplementary material**

Table S1 – Number of quality filtered Illumina® barcoded sequences.

**Figures**

Figure 1 – Experimental design

Figure 2 - Respirometry analysis of Trindade Island coastal soil microcosms. Average CO$_2$ emission rates evaluated during 38 days of incubation. Emissions until 24 days represent the acclimatizing period without oil addition. Readings after 24 days show the differences in CO$_2$ emissions after establishing the two treatments (Control and Oil). The microcosms were incubated at 26 °C and CO$_2$ emission was monitored by an automated respirometer coupled to an infrared CO$_2$ detector.

Figure 3 - Average alpha diversity rarefaction plot for (A) Bacteria/Archaea and (B) Fungi. It shows the number of observed species at a random pool of sequences in different depths.
Figure 4 - Principal coordinate analysis (PCoA). A total of 6 soil samples were analysed by amplicon sequencing. Sequences were rarefied at the same sequencing depth and abundance matrixes were generated using taxa tables summarized at the lowest possible taxonomic level, ranging from phylum to specie. (A) 16S rDNA amplicon sequences coordinates analysis, generated with Weighted Unifrac distance matrix, explaining 90.90% of variation. (B) Fungal ITS1 region amplicon sequences coordinate analysis, generated with Bray-Curtis distance matrix, explaining 96.16% of variation.

Figure 5 - Relative abundance of Bacteria and Archaea phyla using 16S rDNA sequences. Samples are disclosed isolated and as an average of each treatment.

Figure 6 - Relative abundance of Fungi phyla using ITS1 region sequences. Samples are disclosed isolated and as an average of each treatment.
Figure 1 – Experimental design scheme
Experimental design representation

1. **Soil sampling at Trindade Island**
   - Kept in icebox until processing

2. **Soil sieving (< 2mm) and storage at 6 °C.**
   - Preparation of soil stocks (crude oil contaminated and control) for microcosms
   - Soil treatments

3. **Microcosm assembly and incubation of 10 g of soil for acclimatization with CO₂ monitoring**
   - Definition of treatments (soil + hexane and soil + crude oil) after 23 days of acclimatization
   - Incubation for further 15 days, with CO₂ monitoring

4. **End of incubation and soil DNA extraction**
   - Control
   - Crude Oil

5. **20 g of soil per flask**

---
Figure 2 - Respirometry analysis of Trindade Island coastal soil microcosms.

Average CO2 emission rates evaluated during 38 days of incubation. Emissions until 24 days represent the acclimatizing period without oil addition. Readings after 24 days show the differences in CO2 emissions after establishing the two treatments (Control and Oil). The microcosms were incubated at 26 °C and CO2 emission was monitored by an automated respirometer coupled to an infrared CO2 detector.
3

Figure 3 - Average alpha diversity rarefaction plot

Average alpha diversity rarefaction plot for (A) Bacteria/Archaea and (B) Fungi. It shows the number of observed species at a random pool of sequences in different depths.
Alpha Diversity Rarefaction

A

Number of Prokaryotic Observed OTUs

Sequences Per Sample

Control
Crude Oil

B

Number of Fungal Observed OTUs

Sequences Per Sample
Figure 4 - Principal coordinate analysis (PCoA).

A total of 6 soil samples were analysed by amplicon sequencing. Sequences were rarefied at the same sequencing depth and abundance matrixes were generated using taxa tables summarized at the lowest possible taxonomic level, ranging from phylum to specie. (A) 16S rDNA amplicon sequences coordinates analysis, generated with Weighted Unifrac distance matrix, explaining 90.90% of variation. (B) Fungal ITS1 region amplicon sequences coordinate analysis, generated with Bray-Curtis distance matrix, explaining 96.16% of variation.
Figure 5 - Relative abundance of Bacteria and Archaea phyla using 16S rDNA sequences.

Samples are disclosed isolated and as an average of each treatment.
Figure 6 - Relative abundance of Fungi phyla using ITS1 region sequences.

Samples are disclosed isolated and as an average of each treatment.
### Table 1 (on next page)

Table 1 - Summary of physicochemical data for surface soil cores (0 - 10 cm) sampled at the northeast coast of Trindade Island - Brazil.

1. Remaining phosphorus (Alvarez et al., 2000).
2. Extracted with Mehlich – 1.
3. Extracted with monocalcium phosphate in acetic acid (HOEFT et al., 1973).
4. Extracted with KCl 1 mol L-1.
5. Walkley and Black method/OM = C.org * 1.724.
Table 1 – Summary of physicochemical parameters for soil cores (0 - 10 cm) sampled at the northeast coast of Trindade Island - Brazil.

| Characteristic | Unit       | Value   |
|----------------|------------|---------|
| pH – H₂O       |            | 5.6     |
| Soil texture   | Sandy Loam |         |
| P-rem (1)      | mg L⁻¹     | 26.5    |
| P (2)          | mg kg⁻¹    | 1290.8  |
| K (2)          | mg kg⁻¹    | 180.33  |
| S (3)          |            | 5.63    |
| Ca⁺²(4)        | cmol c kg⁻¹| 9.84    |
| Mg⁺²(4)        |            | 2.78    |
| OM             |            | 0.64    |
| N              | %          | 0.19    |
| C (5)          |            | 0.37    |

(1) Remaining phosphorus (Alvarez et al., 2000). (2) Extracted with Mehlich – 1. (3) Extracted with monocalcium phosphate in acetic acid (Hoeft et al., 1973). (4) Extracted with KCl 1 mol L⁻¹. (5) Walkley and Black method/OM = C.org * 1.724.
Table 2 – Average (n = 3) alpha diversity comparison between the treatments control and crude oil for bacteria and archaea groups.

*Two-sample parametric t-test.
Table 2 – Average (n = 3) alpha diversity comparison between the treatments control and crude oil for bacteria and archaea groups.

| Metrics                | Control | Std. Err. | Crude Oil | Std. Err. | p-value* |
|------------------------|---------|-----------|-----------|-----------|----------|
| Rarefaction level      | 45690   | -         | 45690     | -         | -        |
| Good's coverage        | 0.987   | 0.001     | 0.989     | 0.002     | -        |
| PD whole tree          | 177.51  | 1.69      | 164.87    | 3.72      | 0.012546 |
| Chao 1                 | 3107.12 | 39.7      | 2796.51   | 149.7     | 0.047083 |
| Observed species       | 2679.23 | 36.9      | 2443.20   | 78.5      | 0.018392 |

*Two-sample parametric t-test.
Table 3 – Average (n = 3) alpha diversity comparison between the treatments control and crude oil for fungi.

*Two-sample parametric t-test.
Table 3 – Average (n = 3) alpha diversity comparison between the treatments control and crude oil for fungi.

| Metrics              | Control | Std. Err. | Crude Oil  | Std. Err. | p-value* |
|----------------------|---------|-----------|------------|-----------|----------|
| Rarefaction level    | 25315   | -         | 25315      | -         | -        |
| Good's coverage      | 0.99    | 0.001     | 0.99       | 0.001     | -        |
| Chao 1               | 100.58  | 12.1      | 69.96      | 12.9      | 0.0548   |
| Observed species     | 96.46   | 11.8      | 67.8       | 11.7      | 0.0681   |

*Two-sample parametric t-test.
Table 4 – Bacterial/Archaeal OTUs presenting an average absolute abundance significantly different between the treatments “Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

*p-values corrected by the FDR method.*
Table 4 – Bacterial/Archaeal OTUs presenting an average absolute abundance significantly different between the treatments “Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

| Taxa                                      | Control | Crude Oil | q-value*   |
|-------------------------------------------|---------|-----------|------------|
| g__Candidatus Nitrososphaera              | 10275.0 | 4098.0    | 3.77E-03   |
| p__Acidobacteria;o__DS-18                 | 546.0   | 740.0     | 9.05E-03   |
| p__Acidobacteria;o__Sva0725               | 370.0   | 704.0     | 9.20E-05   |
| p__Actinobacteria;g__Iamia                | 93.0    | 206.0     | 1.41E-02   |
| p__Actinobacteria;f__Actinosynnemataceae  | 17.0    | 71.0      | 6.66E-07   |
| p__Actinobacteria;g__Gordonia             | 1.0     | 51.0      | 2.69E-02   |
| p__Actinobacteria;f__Intrasporangiaceae   | 42.0    | 186.0     | 1.03E-07   |
| p__Actinobacteria;g__Nocardioides         | 57.0    | 105.0     | 1.13E-02   |
| p__Actinobacteria;g__Pimelobacter         | 6.0     | 799.0     | 1.82E-19   |
| p__Actinobacteria;g__Amycolatopsis        | 2.0     | 88.0      | 5.19E-10   |
| p__Actinobacteria;f__Streptomycesetaceae  | 166.0   | 3743.0    | 2.19E-48   |
| p__Actinobacteria;g__Streptomyces         | 380.0   | 613.0     | 6.03E-04   |
| p__Actinobacteria;g__Actinomadura         | 32.0    | 71.0      | 3.75E-03   |
| p__Actinobacteria;c__MB-A2-108             | 80.0    | 130.0     | 2.16E-02   |
| p__Actinobacteria;f__Rubrobacteraceae     | 4085.0  | 1590.0    | 1.72E-04   |
| p__Actinobacteria;g__Rubrobacter          | 6674.0  | 1710.0    | 4.18E-08   |
| p__Bacteroidetes;g__Croccinitomix         | 62.0    | 0.0       | 5.30E-11   |
| p__Bacteroidetes;g__Fluvuicola            | 568.0   | 53.0      | 9.23E-05   |
| p__Firmicutes;o__Bacillales              | 109.0   | 33.0      | 3.84E-02   |
| p__Firmicutes;g__Alicyclobacillus         | 102.0   | 30.0      | 1.45E-02   |
| p__Firmicutes;g__Bacillus                 | 1680.0  | 504.0     | 5.32E-05   |
| p__Firmicutes;g__Virgibacillus            | 162.0   | 65.0      | 1.61E-02   |
| p__Firmicutes;g__Cohnella                 | 54.0    | 14.0      | 1.71E-02   |
| p__Firmicutes;f__Thermoactinomycetaceae  | 52.0    | 7.0       | 1.08E-04   |
| p__Nitrospira;g__Nitrospira               | 1356.0  | 591.0     | 8.71E-04   |
| p__Planctomycetes;c__Pla3                 | 169.0   | 65.0      | 4.17E-02   |
| p__Planctomycetes;o__B97                 | 127.0   | 52.0      | 4.41E-02   |
| p__Proteobacteria;c__Alphaproteobacteria  | 95.0    | 214.0     | 1.33E-05   |
| p__Proteobacteria;f__Caulobacteraceae     | 62.0    | 105.0     | 1.02E-02   |
| p__Proteobacteria;g__Phenyllobacterium    | 52.0    | 159.0     | 2.03E-06   |
| p__Proteobacteria;o__Ellin329             | 579.0   | 754.0     | 2.80E-02   |
| p__Proteobacteria;f__Rhizobiaceae         | 142.0   | 223.0     | 1.29E-02   |
| p__Proteobacteria;o__Rhodospirillales     | 1862.0  | 1003.0    | 2.52E-02   |
| Phylum          | Genus            | % of Total Reads | Average Reads Per Sample | p-value   |
|-----------------|------------------|------------------|--------------------------|-----------|
| p__Proteobacteria;g__Phaeospirillum |                | 33.0             | 83.0                     | 6.83E-04  |
| p__Proteobacteria;o__Rickettsiales |              | 55.0             | 13.0                     | 3.75E-03  |
| p__Proteobacteria;f__Alcaligenaceae |              | 88.0             | 152.0                    | 1.25E-02  |
| p__Proteobacteria;f__Burkholderiaceae |           | 1.0              | 110.0                    | 9.60E-14  |
| p__Proteobacteria;g__Burkholderia |                 | 6.0              | 108.0                    | 8.14E-16  |
| p__Proteobacteria;f__Comamonadaceae |            | 611.0            | 4498.0                   | 1.14E-14  |
| p__Proteobacteria;g__Delftia |                     | 15.0             | 289.0                    | 1.34E-09  |
| p__Proteobacteria;g__Cupriavidus |               | 25.0             | 258.0                    | 1.09E-17  |
| p__Proteobacteria;f__Entotheonellaceae |         | 1030.0           | 472.0                    | 1.11E-03  |
| p__Proteobacteria;f__Bacteriovoracaceae |         | 185.0            | 35.0                     | 2.80E-02  |
| p__Proteobacteria;f__Syntrophobacteraceae |     | 7112.0           | 3975.0                   | 3.01E-02  |
| p__Proteobacteria;f__Alteromonadaceae |          | 969.0            | 54.0                     | 5.43E-05  |
| p__Proteobacteria;g__Cellvibrio |                    | 101.0            | 22.0                     | 1.37E-04  |
| p__Proteobacteria;f__Moraxellaceae |                | 3.0              | 271.0                    | 4.08E-02  |
| p__Proteobacteria;g__Acinetobacter |               | 6.0              | 436.0                    | 3.44E-06  |
| p__Proteobacteria;g__Perlucidibaca |                  | 9.0              | 1496.0                   | 3.11E-07  |
| p__Proteobacteria;g__Arenimonas |                    | 60.0             | 9.0                      | 2.93E-04  |
| p__TM7;c__SC3 |                          | 127.0             | 17.0                     | 2.41E-08  |
| p__TM7;c__TM7-1 |                        | 50.0             | 7.0                      | 1.77E-04  |

*p-values corrected by the FDR method.
**Table 5** (on next page)

Table 5 - Fungal OTUs presenting an average absolute abundance significantly different between the treatments “Crude Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

*p-values corrected by the FDR method.
Table 5 – Fungal OTUs presenting an average absolute abundance significantly different between the treatments “Crude Oil” and “Control”, under the EdgeR Fisher’s exact test and $\alpha = 0.05$.

| Taxa                                                       | Control | Crude Oil | q-value*   |
|------------------------------------------------------------|---------|-----------|------------|
| p__Ascomycota;f__Clavicipitaceae                           | 96.3    | 2442.6    | 1.96E-16   |
| p__Ascomycota;g__Fusarium                                  | 834.0   | 19800.6   | 7.25E-16   |
| p__Zygomycota;g__Mortierella                              | 11430.0 | 69846.3   | 2.55E-08   |
| p__Ascomycota;o__Hypocreales;                             | 378.0   | 1148.3    | 1.01E-05   |
| p__Ascomycota;g__Lecanicillium                             | 0.0     | 135.0     | 3.02E-05   |
| p__Ascomycota;f__Bionectriaceae                            | 14704.3 | 202.3     | 8.01E-04   |

*p-values corrected by the FDR method.