

1 **Comparative mesocosm study of biostimulation efficiency in two different oil-amended**
2 **sub-Antarctic soils**

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1 **Abstract**

2 Biological treatment has become increasingly popular as a remediation method for soils and
3 groundwater contaminated with petroleum hydrocarbon, chlorinated solvents, and pesticides.
4 Bioremediation has been considered for application in cold regions such as Arctic and sub-
5 Arctic climates and Antarctica. Studies to date suggest that indigenous microbes suitable for
6 bioremediation exist in soils in these regions. This paper reports on two case studies at the sub-
7 Antarctic Kerguelen Island, in which indigenous bacteria were found that were capable of
8 mineralizing petroleum hydrocarbons in soil contaminated with crude oil and diesel fuel. All
9 results demonstrate a serious influence of the soil properties on the biostimulation efficiency.
10 Both temperature elevation and fertilizer addition have a more significant impact on the
11 microbial assemblages in the mineral soil than in the organic one. Analysis of the
12 hydrocarbons remaining at the end of the experiments confirmed the bacterial observations.
13 Optimum temperature seems to be around 10°C in organic soil while it was higher in mineral
14 soil. The benefit of adding nutrient was much stronger in mineral than in the organic soil.
15 Overall, this study suggests on the basis of microbiological and physicochemical parameters,
16 that biostimulation treatments were driven by soil properties and that ex-situ bioremediation
17 for treatment of cold contaminated soils will allow greater control over soil temperature, a
18 limiting factor in cold climates.

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20 Key words: sub-Antarctic, soil, biostimulation, crude oil, diesel fuel, oil-degrading bacteria

1 **Introduction**

2 The increasing use of petroleum hydrocarbons in high latitude regions led to a growing
3 probability of major spillages threatening terrestrial and aquatic environments. A number of
4 studies have focused on chronic hydrocarbon contamination near the Antarctic and sub-
5 Antarctic research stations, revealing the presence and persistence of these human-derived
6 contaminants (Cripps and Shears, 1997; Snape et al., 2001; Delille and Pelletier, 2002;
7 Aislabie et al., 2004; Rayner et al., 2007). Of all the different types of contamination reported
8 up to now in Antarctica, petroleum has been identified as the most significant problem to be
9 addressed (Snape et al., 2001). The need for research into hydrocarbon degradation process at
10 low temperatures is therefore without question in order to preserve the vulnerable and extreme
11 Antarctic environments. Many physical, chemical and biological technologies have been
12 developed to remove hydrocarbon pollutants from soils and restore the environmental
13 integrity. However, these techniques are often unfeasible in remote areas where accessibility
14 of heavy equipment and materials is limited or prohibited by the high sensitivity of threatened
15 environments. Bioremediation, including biostimulation and bioaugmentation, has proven to
16 be an effective method for cleaning up residual oil in a variety of environments (Van Hamme
17 et al., 2003) and has been proposed as the only viable management option that can be
18 implemented on a large scale in Antarctic environments (Snape et al., 2001). In many
19 instances, however, the rate of petroleum biodegradation in cold environments is severely
20 limited by temperature fluctuation and the available concentrations of fixed forms of nitrogen
21 and phosphate (Walworth and Reynolds 1995; Coulon et al., 2004; Delille et al., 2007). Low
22 temperature plays a significant role in controlling the nature and extent of microbial
23 hydrocarbon metabolism (Gerdes et al., 2005, Nedwell, 1999) and directly affects both the rate
24 of biodegradation and the physicochemical behaviour of oil hydrocarbons, such as viscosity,
25 diffusion and volatilization (Enell *et al.*, 2005; Maliszewska-Kordybach, 2005). Additionally,

1 nutrient supplementation (so-called biostimulation) of polluted sites has been shown to be an
2 effective means of stimulating hydrocarbon biodegradation activities of indigenous microbial
3 populations and thereby of reducing the ecological impact of oil pollution (Delille et al., 2004;
4 Aislabie et al, 2006). However, monitoring studies conducted in oil-contaminated polar soils
5 and experimental research during the last decades have documented the complexity of cold
6 ecology and the parameters that affect the severity of impacts to these systems (see for review
7 Aislabie et al., 2006). This knowledge complicates decisions regarding cleanup in cold
8 environments, because parameters such as substrate type, oil type, season of impact and
9 climate may all affect the eventual recovery of oil-contaminated polar soils. Limiting factors
10 need to be overcome if microbial breakdown of contaminants is to be used effectively
11 (Alexander 1999, Margesin et al., 2007). However, the conditions when cleanup is desirable in
12 a polar soil are not clearly delineated yet, and several important questions remain unresolved.
13 Among them two of the more decisive are: which methods should be employed and at what
14 point intervention is no longer useful? The aim of the present mesocosm study is to evaluate
15 and compare the benefits of temperature increase and slow release fertilizer addition on
16 biodegradation of diesel fuel and crude oil in two different sub-Antarctic soils.

17

18 **Materials and methods**

19 *Field site, sampling and mesocosm set-up*

20 Two soils were collected from areas with no past exposure to hydrocarbon contamination.
21 Both areas were located near the scientific research station “Port aux Français” in Kerguelen
22 Archipelago (49°21’S, 70°13’E). Soil samples were collected from the surface to a depth of
23 about 0.2 m in approximately 20 m² areas. The first selected soil was an organic soil
24 supporting an abundant vegetal cover (*Acaena magellanica*), while the second one was a
25 mineral soil, completely dry and desert. The general physical, chemical and biological

1 properties of the two sub-Antarctic soils have been previously described (Coulon et al., 2004).
2 Mesocosm experiments were conducted in polyethylene containers of dimensions 27 x 24 x 13
3 cm. Plant residues have been removed for the organic soil before use. Soils have been aerated
4 and homogenized before placing 5 kg ($^w/w$) of soil in each mesocosm. The artificially
5 contaminated soils were prepared by direct application of either 100 ml Arabian light crude oil
6 or diesel fuel (initial contaminant concentration = 30 mg g⁻¹ soil dry mass). The composition
7 of Arabian light crude oil and diesel fuel was 52 and 71% of saturated linear and cyclic
8 alkanes, 45 and 28% of aromatics (2 to 5 rings PAH) and 3 and 1% of polar compounds,
9 respectively. The slow release fertilizer used was Inipol EAP 22[®] (CECA S.A., France, C: N:
10 P = 62:7.4:0.7), which is a stable microemulsion consisting of a urea core (the nitrogen
11 source) surrounded by oleic acid carrier, lauryl-phosphate (surfactant and the source of
12 phosphorus), and butoxyethanol (as viscosity reducer). The initial concentration of the
13 nutrients obtained were 1.2 mg N and 0.1 mg P g⁻¹ soil dry mass. For each temperature of
14 incubation (4°C, 10°C and 20°C), six conditions were used: control, Inipol (50 ml), crude oil
15 (100 ml), crude oil (100 ml) + Inipol (50 ml), diesel (100 ml), and diesel (100 ml) + Inipol (50
16 ml). The mesocosms were incubated in the dark under aerobic conditions during 2 months
17 period. They were homogenized twice a month. Samples for chemistry analysis were collected
18 in the surface layer of the soil and were stored at - 20°C until analysis.

19

20 *Bacteriological counts*

21 The changes in bacterial community abundance, comprising total, heterotrophic and
22 hydrocarbon-degrading microorganisms were studied during a 42 days period after
23 contaminant addition. Sampling dates were 7, 15, 30, and 42 days. Triplicate samples were
24 aseptically collected in the surface layer of the soil (from surface to 2 cm under the surface).
25 Total bacteria were determined by acridine orange direct count (AODC) on black nuclepore

1 filters (0.2 μm) using an Olympus BHA epifluorescence microscope according to the method
2 of Hobbie *et al.* (1977). A minimum of 500 fluorescing cells with a clear outline and definite
3 cell shape cells were counted under oil immersion (x 1000) in a minimum of 10 randomly
4 chosen fields. The standard deviation calculated from 3 replicates was found $\leq 15\%$.
5 Heterotrophic microorganisms in each soil sample was made using the spread plate technique
6 on Nutrient Agar 2216 (Oppenheimer and ZoBell 1982), using distilled water in place of
7 seawater. Inoculated plates (six replicates) were incubated for 10 days at 15°C. Hydrocarbon-
8 degrading microorganisms were determined by the most probable number (MPN) method
9 using tubes containing 9 ml of a basal mineral medium (NH_4Cl : 2.0 g L^{-1} , KH_2PO_4 : 0.89 g L^{-1} ,
10 NA_2HPO_4 : 1.25 g L^{-1} , FeCl_3 : 0.6 mg L^{-1}), supplemented with 0.2 ml of Arabian light crude oil
11 and 1 mg L^{-1} resazurin indicator (Mills *et al.*, 1978). After inoculation (3 tubes per dilution),
12 the tubes were incubated at 12°C for 30 days. The standard deviation calculated from 3
13 replicates was found $\leq 20\%$ for both CFU and MPN estimations.

14

15 *Hydrocarbon extraction and analysis*

16 Before extraction, soil samples were freeze-dried and homogenized by screening through a 1-
17 mm sieve. Extraction procedure and gas chromatography-mass spectrometry (GCMS)
18 analysis setting have been previously described (Coulon and Delille, 2006). Each dry soil
19 sample was extracted with a mix hexane:dichloromethane (1:1) overnight and the solution
20 was evaporated under a mild stream of nitrogen on an ice bath to minimize losses of light
21 alkanes and PAH. Very light hydrocarbons such as benzene, pentane and hexane could be
22 partly lost during freeze drying. These light hydrocarbons are not accounted in total alkanes
23 (calculated from C10 to C36 linear alkanes and including iso-alkanes) and total aromatics
24 (from naphthalene to 4-ring aromatics including alkyl-substituted compounds). Analytes of
25 Arabian light crude oil were normalized to the conservative biomarker 17 α (H), 21 β (H) C30-

1 hopane naturally present in crude oil (Butler *et al.*, 1991) while analytes of diesel fuel were
2 normalized to chrysene, considered to be one of aromatics most resistant to biodegradation
3 (Bossert and Bartha, 1986). Hopane was absent from diesel fuel samples. For quality control,
4 a 1.0 ng μl^{-1} diesel standard solution (ASTM C₁₂-C₆₀ quantitative, Supelco) and a 1.0 ng μl^{-1}
5 PAH Mix Standard solution (Supelco) were analyzed every 20 samples. The variation of the
6 reproducibility of the extraction and quantification protocol was determined by successive
7 extractions and injections of 8 replicates of the same field sample and estimated to be $\pm 10\%$.

8

9 *Statistical analysis*

10 Measures analysis of variance (ANOVA) in Statview F-5.0 PPC (SAS Institute Inc.) was used
11 to analyze the response variables (CFU, MPN, fertilizer and oil analytes) in the two soils
12 separately. When the ANOVA indicated significant differences ($p < 0.05$), univariate
13 ANOVAs were run on data at each time point. Where significant differences were indicated at
14 a specific time point ($p < 0.05$), protected least significant difference (LSD) mean separations
15 were used to assess treatment differences

16

17 **Results**

18 There were only slight changes in total microbial abundance after contamination (data not
19 shown). With values comprised between 10^8 and 10^9 cells g^{-1} , the total number of micro-
20 organisms was roughly constant throughout all the mesocosm experiments.

21 The basal level of heterotrophic microbes in the mesocosms, prior to oil addition, was
22 estimated to be 3.2×10^6 CFU g^{-1} in organic soil and 2.6×10^5 CFU g^{-1} in mineral soil. These
23 number remained relatively constant in most of the uncontaminated mesocosms. Following
24 diesel fuel (Figure 1) or crude oil (Figure 2) addition, the number of heterotrophic microbes
25 did not change significantly in both soils ($P = 0.659$ in organic soil and $P = 0.317$ in mineral

1 soil). Inipol amendment did not induce significant enhancement of the heterotrophic
2 assemblage in organic soil. In contrast significant increases of the number of saprophytic
3 bacteria occurred in all Inipol amended mesocosms of mineral soil ($P < 0.001$). Enhancements
4 reach three orders of magnitude and seems relatively independent from temperature.

5 Before contamination, initial level of oil-degrading microbes estimated by the most probable
6 number (MPN) procedure was 2.7×10^5 MPN g^{-1} in organic soil and 3.0×10^5 MPN g^{-1} in
7 mineral soil (Figures 3 and 4). After crude oil amendment, the number of oil-degrading
8 microbes increased by one order of magnitude in both kind of soils ($P < 0.082$ in organic soil
9 and $P < 0.056$ in mineral soil) over a period of 7 days. In contrast, these numbers did not
10 increase after diesel fuel amendment, except in organic mesocosms incubated at 4 and 20°C.

11 According to the type of oil used, the comparison of the oil-degrading microbe's number for
12 the same soil did not revealed significant difference between diesel fuel and crude oil ($P >$
13 0.05). The efficiency of the biostimulation treatments was much higher in the mineral soil
14 (more than three orders of magnitude) than in the organic one (less than two orders of
15 magnitude) for both kinds of oil contaminants. At the end of the experiment, the numbers of
16 oil-degrading microbes in diesel amended mineral soil microcosms can be four orders of
17 magnitude higher than those determined in corresponding pristine soil whilst this difference is
18 less than two orders of magnitude in organic soil. Inipol amendment alone induced also a
19 significant increase in numbers of oil-degrading microbes in all mesocosms. This increase was
20 significantly greater in mineral soil mesocosms ($P < 0.001$) than in organic ones ($P = 0.056$).

21 Temperature elevation had a slight but positive effect on biostimulation of oil degrading
22 microbes. With the exeption of diesel contaminated organic soil, the maximum numbers were
23 always observed in mesocosms incubated at 20°C.

24 Overall, the total extractable petroleum hydrocarbons content (TPH) of both oil contaminants
25 have reduced by more than 70% and 80% in organic soil mesocosms and more than 76% and

1 96% in mineral soil mesocosms incubated at 4 and 20°C, respectively. Detailed changes of oil
2 hydrocarbon fraction of both crude oil and diesel fuel are shown in Table 1. After fertiliser
3 application, the differences in extent of degradation were most pronounced for both aliphatic
4 (P < 0.001) and aromatic (P = 0.098) fractions in mineral and organic soil mesocosms (Table
5 1; Fig 5). However, the benefit of adding nutrient was higher in mineral than in organic soil.
6 The mass fraction of aromatics relative to aliphatic hydrocarbons has increased of more than
7 20% in crude oil amended mesocosms and more than 24% in diesel fuel amended ones,
8 regardless the soil type. These mass balance shifts were even higher in fertilised mineral soil
9 mesocosms: more than 35% in crude oil amended mesocosms (P < 0.001) and more than 42%
10 in diesel fuel amended ones (P < 0.05). The benefit of increasing temperature is particularly
11 obvious in diesel contaminated mineral soil mesocosms. In contrast, the best results of oil
12 hydrocarbons degradation were observed in organic soil mesocosms incubated at 10°C.

13

14 **Discussion**

15 The present observations demonstrate the clear stimulating effect of oil addition on indigenous
16 bacteria in sub-Antarctic soil. Several orders of magnitude increase in bacterial abundance
17 occurred after both diesel and crude oil addition. These observations are in good agreement
18 with previous results obtained from *in situ* studies in sub-Antarctic soils (Crozet Island)
19 (Coulon & Delille 2006, Delille et al., 2001, 2004). Biostimulation induced a clear increase of
20 the number of hydrocarbon-degrading microbes. As noted by Rivet et al. (1993), some
21 increases in bacterial numbers after Inipol EAP 22 addition may be attributed to the bacteria
22 growing on the oleic acid in the fertilizer. However the concomitant reduction of the residence
23 time of the contaminants demonstrates the efficiency of the biostimulation. Despite that the
24 same general pattern was observed in the two soils, the results indicate a serious influence of
25 the soil properties on the observed biodegradation efficiencies.

1 The intensity of hydrocarbon biodegradation in soil is influenced by a number of site-specific
2 factors (e.g. low temperature, low nutrient availability, low oxygen levels, soil structure,
3 etc...). Among them, it is well established that nutrients are one of the major factors limiting
4 hydrocarbon metabolization in soils (Mohn and Stewart, 2000). Inputs of large quantities of
5 carbon sources (i.e., hydrocarbon contamination) tend to result in rapid depletion of the
6 available pools of major inorganic nutrients, such as nitrogen and phosphorus. Several studies
7 have reported favourable effects of fertilizers on oil biodegradation at low temperatures in
8 Arctic (Braddock *et al.*, 1997, Whyte *et al.*, 1999; Mohn and Stewart, 2000), alpine (Margesin
9 and Schinner 1997a,b; Margesin and Schinner 1999, Margesin *et al.* 2007) and Antarctic soils
10 (Kerry 1993; Wardell 1995; Aislabie *et al.*, 1998; Delille 2000, Powell *et al.*, 2006).

11 A possible reason for the inability of Inipol EAP 22[®] to greatly enhance hydrocarbon-
12 degrading microbes growth in organic soil is that nitrogen and phosphorus are not the major
13 limiting factors in this soil. Initial nitrogen concentrations naturally present were probably
14 high enough to sustain rapid intrinsic rates of biodegradation without addition of fertiliser. In
15 addition, particle size distribution, elemental analysis and water content of both sub-Antarctic
16 soils were strongly different from each other. Thus, differences in the observed degradation
17 rates in both studied soils may be associated with the availability of the contaminants.
18 Furthermore, long-term in situ experiments demonstrated that the wet organic soil seems to be
19 more efficient to retain some toxic compounds than the mineral one (Delille *et al.* 2007). This
20 difference could have a direct influence on the mineralisation rate of hydrocarbons.

21 Microbial metabolism is usually considered as a direct function of the temperature of the
22 environment (Leahy and Colwell 1990). Results of the present mesocosm experiments
23 indicate that a temperature increase can stimulate the microbial hydrocarbon degradation in
24 sub-Antarctic soils. However, the influence of temperature can differ greatly from a soil to
25 another. We have no clear explanation of the difference of temperature sensitivity observed

1 between the two soils used in the present experiment. However, these differences exist and
2 cannot be forgotten.

3 In conclusion, the results obtained under the same experimental design differ greatly
4 from one soil to another, demonstrating a serious influence of the soil properties on the
5 biostimulation efficiency. While spill management requires the development of a quick action
6 in response planning, the finding of this study reinforces the need to evaluate firstly the
7 factors that control both the microbial activity and the degradation of organic compounds at a
8 specific site. Whilst acknowledging the fundamental limitations of ex-situ bioremediation
9 approach in cold environments, this study demonstrated that ex-situ bioremediation is likely to
10 be the strategy of choice for remediation of hydrocarbon-contaminated sub-Antarctic soils.

11

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Legends

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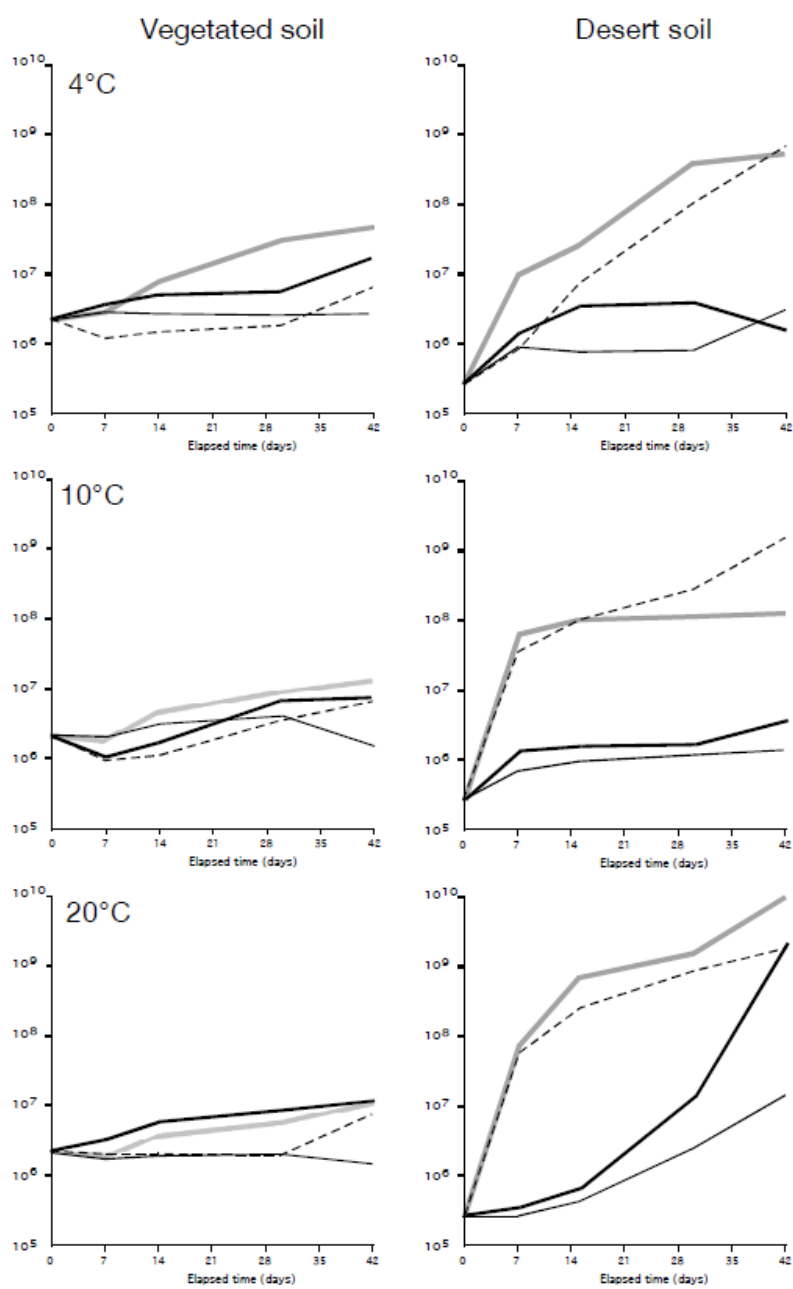
Figure 1: Changes of heterotrophic bacterial abundance during incubation of the diesel amended mesocosms (thin line: control, dotted line: control + fertiliser, bold line: diesel, gray line: diesel + fertiliser).

Figure 2: Changes of heterotrophic bacterial abundance during incubation of the crude oil contaminated mesocosms (thin line: control, dotted line: control + fertiliser, bold line: crude oil, gray line: crude oil + fertiliser).

Figure 3: Changes of hydrocarbon-degrading bacterial abundance during incubation of the diesel contaminated mesocosms (thin line: control, dotted line: control + fertiliser, bold line: diesel, gray line: diesel + fertiliser).

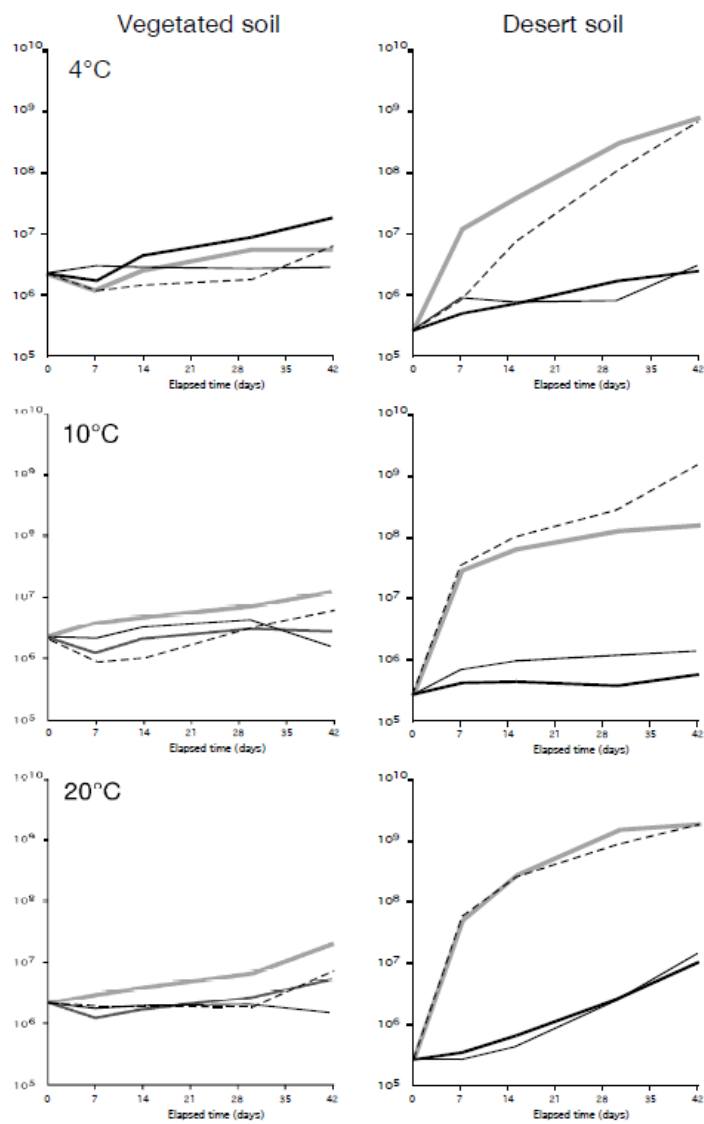
Figure 4: Changes of hydrocarbon-degrading bacterial abundance during incubation of the crude oil contaminated mesocosms (thin line: control, dotted line: control + fertiliser, bold line: crude oil, gray line: crude oil + fertiliser).

Figure 5: Changes in oil hydrocarbon fractions concentration (as % of initial values) over 180 days. Each percentage represents the mean of hydrocarbons fractions from duplicate samples and bars indicate standard deviation.



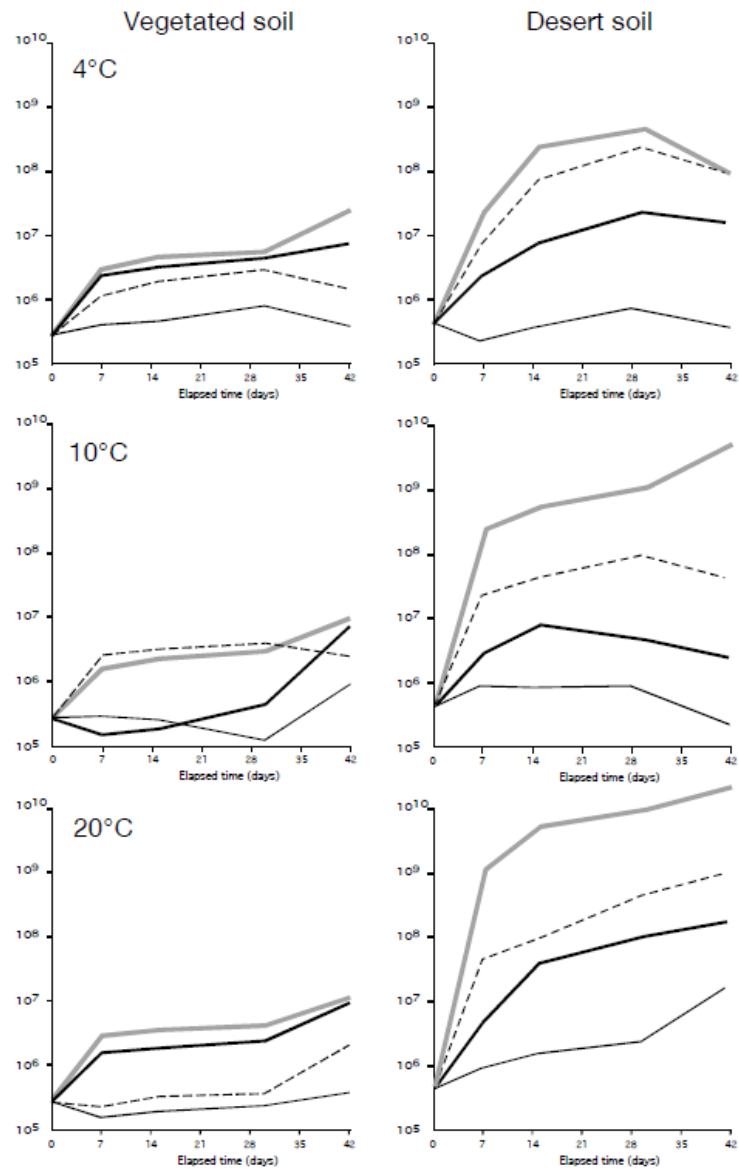
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2 Figure 1



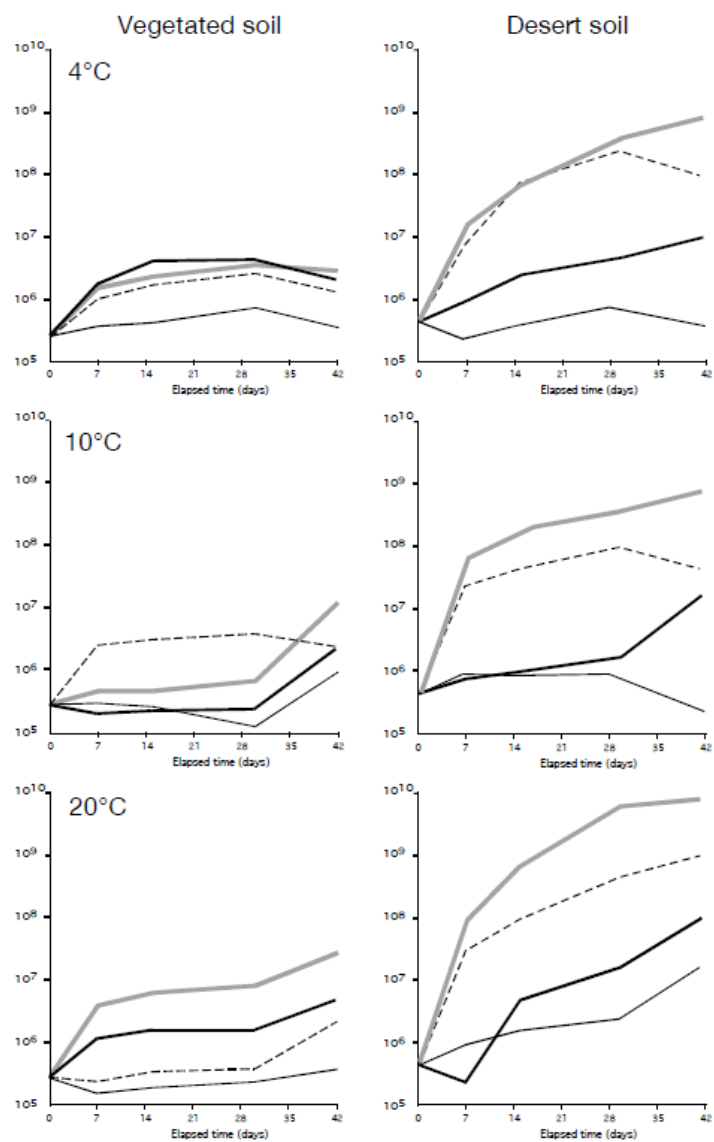
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2 Figure 2



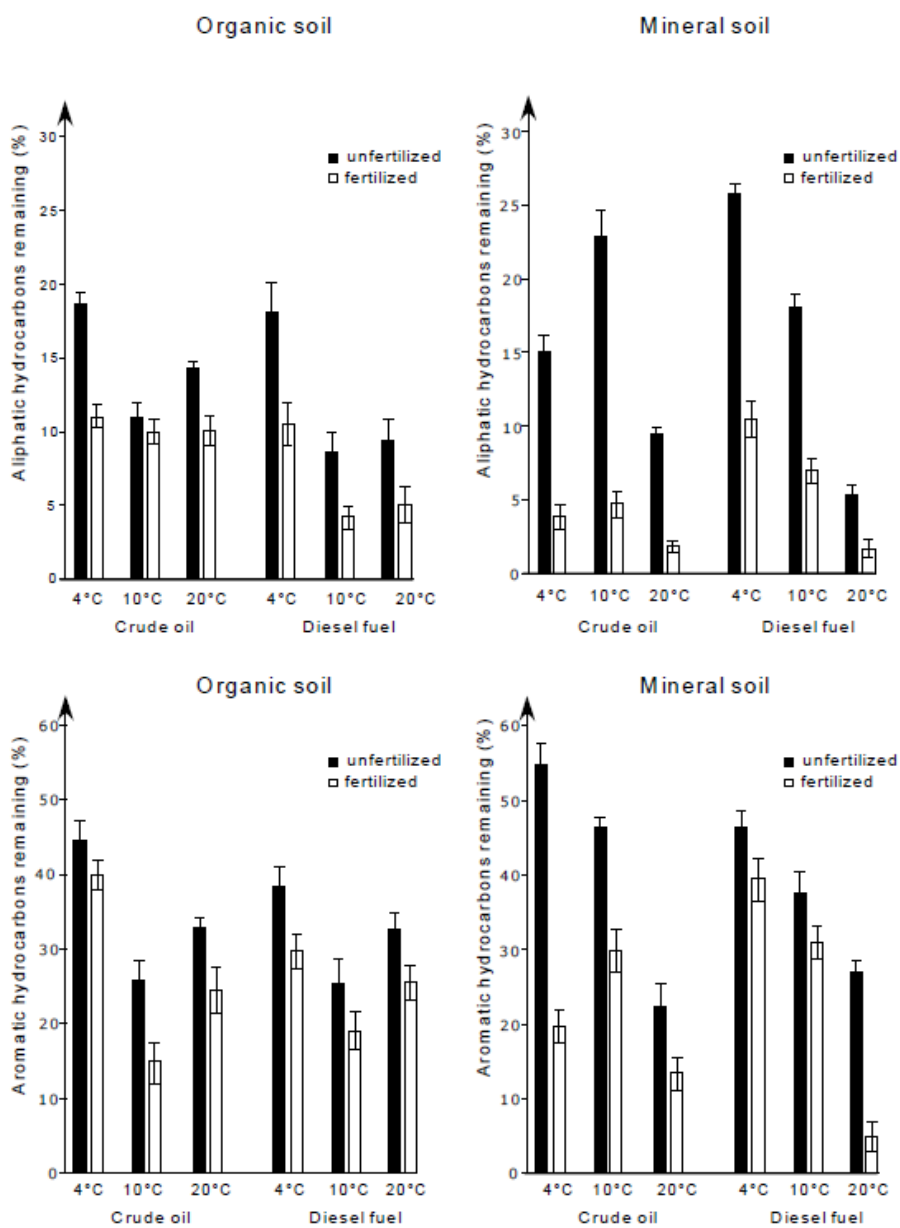
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2 Figure 3



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2 Figure 4



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2 Figure 5

1 **Table 1:** Changes in oil hydrocarbon fractions concentration (as % degradation of initial
 2 value) over 180 days. Each percentage represents the mean of hydrocarbons fractions from
 3 duplicate samples
 4
 5

Oil type/ treatment	Hydrocarbon fraction	Organic soil			Mineral soil		
		4°C	10°C	20°C	4°C	10°C	20°C
Crude oil	<i>Aliphatic</i>						
	EC >10 - 12	65	82	73	82	75	91
	EC > 12 -16	59	79	75	73	70	84
	EC > 16 -35	45	56	53	67	61	71
	<i>Aromatic</i>						
	EC >10 - 12	63	65	77	76	79	87
	EC > 12 -16	55	69	62	54	70	89
	EC > 16 -21	23	39	42	38	46	51
	Crude oil + inipol	<i>Aliphatic</i>					
EC >10 - 12		81	84	89	96	96	> 99
EC > 12 -16		76	79	85	90	95	98
EC > 16 -35		67	53	53	86	90	95
<i>Aromatic</i>							
EC >10 - 12		73	85	77	80	87	89
EC > 12 -16		55	69	62	70	74	79
EC > 16 -21		39	42	49	65	69	72
Diesel		<i>Aliphatic</i>					
	EC >10 - 12	82	94	98	79	90	91
	EC > 12 -16	69	77	89	75	89	84
	EC > 16 -35	45	51	53	58	66	71
	<i>Aromatic</i>						
	EC >10 - 12	67	74	67	73	81	88
	EC > 12 -16	58	61	59	65	69	77
	EC > 16 -21	39	45	42	43	56	63
	Diesel + Inipol	<i>Aliphatic</i>					
EC >10 - 12		85	98	> 99	94	> 99	> 99
EC > 12 -16		78	86	89	88	97	98
EC > 16 -35		56	71	69	67	71	82
<i>Aromatic</i>							
EC >10 - 12		75	82	76	84	89	94
EC > 12 -16		63	71	62	68	71	85
EC > 16 -21		51	65	59	49	58	71