Bioremediation for a Soil Contaminated with Hydrocarbons

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Abstract

The objective was to remediate, through biopiles, 27,400 m³ of soil contaminated with heavy fraction hydrocarbons (HFH) at a Maritime Terminal in southeastern Mexico. To clean the soil, two bioremediation stages were considered: i) batches of soil of 3,800 m³ subjected to land farming pretreatment, ii) biopiles formation of 70 m length, 16 m width, and 2 m height. The parameters controlled during 8 months of bioremediation were: nutrients, water content, aeration, and temperature. Monitoring contemplated monthly sampling of each biopile, the analyzed parameters were the heavy fraction hydrocarbons, bacteria counting, and respirometry. Results revealed that HFH removal in biopiles reached 43.2% after pre-treatment (land farming), 68.7% at mid-treatment, and 77.7% at the end of treatment. The land farming pretreatment achieved a higher percentage of hydrocarbons removal, because aeration was very efficient at the start of treatment; therefore, enhancing aerobic biodegradation.

Keywords: Bioremediation; Biopiles; Land farming; Contaminated soil; Hydrocarbons degradation; Heavy fraction hydrocarbons

Introduction

Bioremediation of soils is currently a relevant issue, because it implies a process in which organic contaminants in the subsoil are biodegraded and can become mineralized so that eventually they become non-toxic compounds. The contaminant does not enter another physical state because it is degraded.

Bioremediation is aimed at maintaining the maximal possible growth of microorganisms until the carbon source (organic contaminant) decreases and, consequently, the microbial population decreases [1].

Physical and chemical factors are needed for an efficient bioremediation process; including water, temperature, pH, oxygen, and major and minor nutrients.

- **Water**: Water content is one of the most important factors for degradation, since water constitutes 80 to 90% of the weight in the molecular composition of bacterial cells and is the main nutrient [2].

- **Ph**: The intracellular pH value lies between 6.5 and 7.5, hence this is the required pH range needed for optimal microorganisms’ growth.

- **Temperature**: The chemical and enzymatic reactions of the cell increase concomitantly with increasing temperature. There are: a minimal temperature for each organism, below which no growth occurs, an optimal temperature at which growth is faster, and a maximal temperature above which no more growth occurs. The temperature range considered optimal for heterotrophic aerobic bacteria is between 20°C and 35°C [3].

- **Oxygen**: Oxygen is the electron acceptor most used by microorganisms to degrade organic compounds in an aerobic environment. If the oxygen content of the soil is below 2 mg/L, conditions are favorable for an anaerobic environment.

- **Nutrients**: The solid portion of the bacterial cell is constituted by carbon, nitrogen, hydrogen, phosphorus, and, to a smaller extent, potassium, calcium, magnesium, chlorides, iron, and others. The main component (50%) is carbon. The contaminant to be degraded must contain this element. Oxygen, with 20%, is the second most abundant element in the cell. Oxygen is needed for new cells and as electron acceptor, hence, it is necessary to count upon large amounts of oxygen for biological degradation. The other major nutrients required by microbiorganisms are nitrogen and phosphorus. The three main nitrogen sources in microbiorganisms are proteins, cell wall constituents, and nucleic acids. Phosphorus, in the form of phosphates, is used by microbiorganisms to synthesize phospholipids and nucleic acids [4].

Factors that might limit the activity of microbiorganisms are low temperatures, very low or very high pH values, chemical agents, such as heavy metals, halogens, organic and oxidizing contaminants.

The main techniques applied to bioremediation are: in situ bioremediation, biopiles, land farming, phytoremediation, bioaugmentation, bioventing.

The biopiles technique consists in forming piles with the contaminated soil and stimulating the microbial communities through aeration and/or by adding nutrients and water. The increment in microbial activity is directly proportional to the reduction in heavy fraction hydrocarbon (HFH) concentrations. Biopiles are aimed at reducing the concentration of hydrocarbons that are adsorbed in contaminated soils by means of biodegradation [5]. Biopiles is the most commonly used technique to treat soils contaminated with petroleum hydrocarbons, especially soils having a predominantly sandy granulometry [6-8].

Microbiological activity can be stimulated by supplying oxygen, through aeration, and water and nutrients, such as nitrogen and phosphorus.

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Efficiency of a biopile depends on several parameters, which are grouped in three categories (Fahnstock et al, 1998), these are:

1. Soil characteristics.
2. Characteristics of the contaminants.
3. Weather conditions.

The type of soil is very important because water, nutrients, and air must be able to migrate with some ease through the soil pores to allow microorganisms to accomplish degradation. Texture of the soil influences soil permeability, water content, and soil density. Highly permeable soils are the most easily aerated and, therefore, are the most adequate to be used for biopiles.

Biopiles are constructed on an impermeable base to reduce the possible migration of lixiviates towards the subsoil. In addition, a network of perforated tubes are installed within the biopile and connected to an aeration system to allow air penetration into the soil and supplying air to bacteria.

Nature of soil contamination

The objective of this work was to clean a soil contaminated with hydrocarbons by means of bioremediation, at a contaminated site located within a PEMEX facility in the state of Tabasco, in southeastern Mexico, where the largest petroleum extraction and production processes of the country are carried out.

Soil contamination was present on the site, predominantly as petroleum hydrocarbons. This contamination resulted from storage and handling of fuels and residual petroleum from storage tanks. The petroleum hydrocarbons were heavier fractions (HFH) (C15-C36) than expected from the nature of the activities on the site, with concentrations of up to 20000 mg/kg.

Two areas of soil contamination were identified requiring remediation. Zone A and Zone B (Figure 1). The site had been contaminated since 40 years ago due to inadequate storage practices. The bioremediation area (Zone C) was located at 300 m from Zone B.

The contaminated site, at the Marine Terminal “Dos Bocas”, in the state of Tabasco, is located in an area of 11 150 m², with an irregular depth between 1 and 6 m, with an average hydrocarbons concentration of 20000 mg/kg, and a contaminated volume of 27 400 m³. Remediation of the soil involved excavation of all the soil exceeding 6000 mg/kg of heavy fraction hydrocarbons, as specified by the Mexican normativity. The water table was found at an average of 3.40 m depth.

Materials and Methods

At the start of the work, the soil was characterized and its main parameters (pH, porosity, organic matter content, soil granulometry, water content, content of heterotrophic bacteria) were determined.

To clean the soil, two bioremediation stages were considered: i) batches of soil were prepared with an approximate volume of 3 800 m³ that were pretreated through land farming. In this stage, the necessary calculated nutrients were mixed and added to the soil; the soil was aerated and mechanically homogenized during 28 days to go over to the second stage. ii) The second stage consisted of forming biopiles of 70 m length, 16 m width, and 2 m height. The general procedure was as follows.

Extraction and separation of the clean soil

The clean soil was separated from the contaminated soil in the study area, starting at the point where contamination was observed. During this activity, it was observed that the distribution of the contamination was very variable, since there were zones in which contamination appeared at 0.7 m depth, whereas, in other zones, the contaminated soil appeared at more than 4.0 m depth. The clean soil was carried over from Zone A to Zone B, where it was maintained until the time was deemed adequate to fill-out the cleaned areas. Separation of the clean soil from the contaminated one was performed based on the organoleptic characterization, and considering the results of the samples sent to a certified laboratory to be analyzed.

Extraction and transportation of the contaminated soil

The contaminated soil was extracted in land strips of approximately 15 m width, with a Caterpillar 320 excavator, until a depth ranging from 5.0 to 6.50 m. The criterion to stop excavation consisted in collecting samples at the excavated depths for analysis and once the analytical results indicated that the HFH values where within the maximal permissible limits (MPL), according to the Mexican normativity; continuing, then, with the next strip of land.

- Cleaned areas: Once reaching the depth at which no contamination was detected, the area was filled with the initially separated clean soil that had been stored at the same site.
- Soil-sieving: Once the contaminated soil had been extracted and separated from the clean soil, it was carried over to Zone C, where the final separation of thick materials existing in the soil was performed by means of a 4 m wide, 3.5 m high, and 8 cm mesh-size soil-sieve.
- Land farming (pretreatment): Once the soil had been sieved, it was taken to the homogenization, aeration, and nutrients addition processes. To optimize soil conditions, the soil was mixed and aerated mechanically, and nutrients were added according to a C:N:P ration of 100:10:1, to promote the biodegradation process of the contaminants.
- Calculations were based on the initial HFH concentration of 20000 mg/kg, considering a carbon concentration of 16000 mg/kg (80%) of HFH and adjusting the effective nutrients content in the agricultural fertilizers used for this purpose.
- Biopiles: Before starting the biopile construction, a high density, 1-mm thick polyurethane liner was placed as bedding to protect the subsoil from an eventual generation and migration of
leachate. Each biopile was 60-m long, 16-m wide, 2-m high, with a 1:1 slope. They were constructed in layers of 40-cm thickness and at the end of each layer, a grooved 1-inch diameter PVC tube was placed along the width of the biopile, at a 3.50-m distance, to provide air inside the biopile.

- **Aeration system**: Oxygen (2 kg) is needed for each kilogram of hydrocarbons. The constructed biopiles had a volume of 1624 m³, each, and a bulk density of 1.6 ton/m³, i.e., a mass of 2598 ton. Hence, each biopile contained 36 372 000 kg of hydrocarbons, that means that a supply of 83 ton of O₂ is needed. It is considered that this amount of air should be supplied in 150 days, at a flow of 553 kg/day = 384 g/min. In a work performed, at an actual scale, in biopiles bioremediation of air should be supplied in 150 days, at a flow of 553 kg/day = 384 g/min. In a work performed, at an actual scale, in biopiles bioremediation of air should be supplied in 150 days, at a flow of 553 kg/day = 384 g/min.

Because of the high temperatures at the site, together with the temperature at the compressors output (50°C), it was necessary to design a cooling system to be placed on the output of the compressor. In this way, the average temperature of the air at the entrance to the biopile was 30°C, which is an adequate temperature for the growth of the microorganisms in charge of degrading the contaminant.

Biopiles were maintained at optimal conditions by means of an aeration system and water supply using constant irrigation. To protect the biopiles, a gravel layer was added at the end of the construction, placed on the surface, to avoid erosion of the biopile’s soil by wind and rain.

### Soil sampling of biopiles

To assess periodically the HC concentrations in each biopile, these were monitored monthly. Twelve sampling points were considered at each biopile, at three different depths in the lateral sides of the biopile (0.5, 1.5, and 2.5 m) and at 0.5, 1.2, and 1.8 m depth in the upper side, as shown in Figure 1. That is, 36 samples were obtained from each biopile; by making a compound sample at each depth, nine samples are obtained for analyses.

### Results and Discussion

Table 1 presents the results of the physical, chemical, and biological parameters of the soil.

#### Removal of heavy fraction hydrocarbons

At the start of the project, once the contaminated soil had been sieved and stored, 10 samples were taken from diverse portions of the stored soil. Two compound samples were obtained and considered as the initial concentration of the contaminated soil. The average concentration obtained was of 20 213 mg/kg.

In another work, bioaugmentation and natural attenuation were tested as remediation strategies in biopiles, sampled periodically until reaching 182 days. Biopiles were divided in quadrants and samples taken from each quadrant, then compound samples from the first samplings were made and sieved to separate the fraction equal or smaller than 2 mm [10]. Other authors tested composting in once constructed; biopiles were sampled as shown in Table 2.

Table 3 shows the average results obtained from the heavy fraction hydrocarbons monitoring. Results obtained in each biopile were averaged. Biopiles for bioremediation of hydrocarbons-contaminated soil. To sample the biopiles that were 2 to 4 m high, a compound sample was made for each biopile constituted by 10 to 12 simple samples, which were then homogenized and sieved to take the sample [11].

**Table 3: Summary of HFH concentrations in biopiles (mg/kg).**

| Biopile | Initial average | After pretreatment | Mid-treatment average | Final average |
|---------|----------------|--------------------|----------------------|---------------|
| 1       | 20 213         | 9078               | SD                   | 3556          |
| 2       | 20 213         | 12 465             | 6434                 | 5171          |
| 3       | 20 213         | 12 465             | 7091                 | 5610          |
| 4       | 20 213         | 12 465             | 4944                 | 3580          |
| 5       | 20 213         | 12 465             | 5302                 | 3579          |
| 6       | 20 213         | 11 393             | 7685                 | 4899          |
| 7       | 20 213         | 11 393             | 7755                 | 5098          |
| 8       | 20 213         | 6780               | 6093                 | 3966          |
| 9       | 20 213         | 12 700             | 7748                 | 5541          |
| 10      | 20 213         | 12 700             | 5313                 | 4588          |
| 11      | 20 213         | 13 660             | 6030                 | 4043          |
| 12      | 20 213         | 11 308             | 5491                 | 5089          |
| 13      | 20 213         | 11 308             | 5995                 | 5410          |
| 14      | 20 213         | 9657               | SD                   | 3032          |

1 Results of the soil sample from the original site, taken by the quartering method
2 Results of the soil sample for each constructed biopile at mid-time of treatment
3 Average results of sampling on the three sides and the three depths of the biopile SD=No data, HFH= Heavy fraction hydrocarbons

**Table 1: Physical, chemical, and biological parameters of the contaminated soil.**

| Parameter                      | Analytical method                          | Result     |
|--------------------------------|--------------------------------------------|------------|
| HFH                            | EPA0071-B for HFH extraction EPA 1684A for chromatographic analysis | Table 3    |
| Bulk density (g/cm³)           | Method AS 03 (NOM-021-RECNAT-2000)         | 1.2451     |
| Dry density (g/cm³)            | ASTM, D854-83                             | 2.6086     |
| Porosity (%)                   | Difference between bulk and dry density    | 0.52       |
| pH                             | ASTM-D 4972-89                            | 7.16       |
| Organic carbon fraction, fOC (%) | Method AS 07 (NOM-021-RECNAT-2000)         | 2.281      |
| Organic matter (%)             | Method AS 07 (NOM-021-RECNAT-2000)         | 3.965      |
| Granulometric analysis         | SUCS (ASTM, D 422)                        | Sand       |
| Count of aerobic bacteria (CFU/g) | NOM-092-SSA1-1994                          | 8×10^8     |

**Table 2: Sampling frequencies of biopiles.**

| Biopile | Initial average | After pretreatment | Mid-treatment average | Final average |
|---------|----------------|--------------------|----------------------|---------------|
| BP1     | Jan Apr        | ---                | ---                  | ---           |
| BP2     | Mar 10 Apr     | ---                | ---                  | ---           |
| BP3     | Mar 10 Apr     | ---                | ---                  | ---           |
| BP4     | Apr 18 May     | ---                | ---                  | ---           |
| BP5     | Apr 19 May     | ---                | ---                  | ---           |
| BP6     | May 15 Jun     | ---                | ---                  | ---           |
| BP7     | May 22 Jun     | ---                | ---                  | ---           |
| BP8     | Jun 27 Jul     | ---                | ---                  | ---           |
| BP9     | Jun 2 Jul      | ---                | ---                  | ---           |
| BP10    | Jun 27 Jul     | ---                | ---                  | ---           |
| BP11    | Jul 8 Jul      | ---                | ---                  | ---           |
| BP12    | Jul 15 Aug     | ---                | ---                  | ---           |
| BP13    | Jul 25 Aug     | ---                | ---                  | ---           |
| BP14    | Aug 10 Dec     | ---                | ---                  | ---           |
treatment, and 77.7% at the end of treatment. It must be noted that HFH concentrations in the biopiles were different because the soil, although coming from the same area, was from different locations and depths, and had different initial HFH concentrations. Removal was reached after 4 to 9 months of soil treatment. This effect has been studied previously, achieving reductions of 2400 mg/kg to 700 mg kg\(^{-1}\) in 5 months, which represents an average of 70% reduction.

It is considered that the reduction of contaminants was through degradation as they are heavy fraction hydrocarbons and contain no volatile compounds, or might have them at a very low proportion. It has been reported that the portion of hydrocarbons (mid and heavy fraction) reduced due to volatilization is, in general, less than 10% at 25°C during the first 30 days [12].

For biopile 1, the first sampling revealed that the HFH concentrations were already below the maximal permissible limits (MPL) indicated by the NOM-138-SEMARNAT/SS-2003 for HFH from industrial-use soil (6000 mg/kg of soil) (DOF, 2005). Therefore, after the third sampling, in which no contaminants were found, the biopile was dismantled and the clean soil was carried over to the original site to be used as filling material. This fast response could have been due to the fact that the soil extracted for this biopile did not have a high HFH concentration at the start and came from the surface of the contaminated site, where the soil is enriched with aerobic microbial populations with a high potential for hydrocarbons degradation and a higher content of nutrients, as well as better aeration; thus, by stimulating the microbial population with optimal conditions, the contaminant was degraded faster.

The maximal degradation of organic contaminants is usually achieved in the first 2 months; this pattern is similar in soils contaminated with both types of hydrocarbons, mid and heavy fraction. This same pattern has been reported by many authors independently from the initial concentration [11].

Other researchers have reported 48% degradation from an initial total petroleum hydrocarbon (TPH) concentration of 10000 mg/kg [13,14]. These were field-scale biopiles with wood chips as bulking agent. Bench-scale experiments from the same site showed a slightly more effective degradation of 80% [15], but still the same pattern.

Biopiles 2 and 3 were the slowest ones in responding to the cleaning treatment, they yielded results below the MPL until the sixth month of treatment; this can have been due to the higher initial concentration and because the soil came from deeper layers, reaching the phreatic level. In this stratum, the particles size is much smaller than at the surface, corresponding to a clayish-lime stratum; hence, oxygen transfer at this level is so low that it does not allow for the survival of aerobic microbial populations. We believe that the bacteria that might have been stimulated were facultative populations with some potential for hydrocarbons degradation, which, when subjected to optimal aerobic condition, were able to develop and degrade the contaminant; however, the developing process was slower than if the organisms had been strictly aerobic. In biopiles with different aeration systems, it has been demonstrated that a high water content has a negative impact on the biodegradation process; hence, more air has to be introduced than that naturally entering the biopiles [16].

In biopiles 4, 5, 6, and 7, results were below the MPL between the third and fourth month of treatment. The behavior was the expected one, because they were within the time range theoretically assumed for a biopile under the conditions of the site described herein. Although the soil could have contained also fine fractions, the speed at which the soil was extracted was higher than the speed at which the biopiles were constructed; hence, the soil was under aerobic conditions for a longer time than biopiles 2 and 3. Thus, it is assumed that the microorganisms had a longer adaptation time, and when the biopiles were constructed after the pretreatment, the developing stage of the microorganisms could have been optimal, and degradation of contaminants was not so slow. Other authors have demonstrated the importance of aeration in this type of treatments, since their experiments with aeration reached a reduction of up to 95% in the first or second month of treatment, as compared to the control without aeration that did not depict a significant reduction in contaminants concentration [17–19].

For biopiles 8, 9, and 11, values near the MPL were obtained in the second month of sampling, but, on the next sampling, they showed an 8 to 10% increase, to finally decrease and reach concentrations below the MPL after 4 months. This increase could be attributed to the fact that heavy rains occurred during pretreatment and construction stages of these biopiles, which could have induced soil-washings, causing a higher availability of hydrocarbons that was reflected in an increase in concentration; however, since the increase in concentration was not too large, it did not cause a negative impact on the microbial population, reaching sufficient removal of hydrocarbons.

For biopiles 10, 12, and 13, the initial concentrations were very close to the MPL, and the removal was slow, but did reach the MPL, this could have been due to the fact that the degradation rate becomes slower with low hydrocarbons concentration.

Biopile 14 was the one to reach concentrations very much below the MPL in the shortest time, as established by the NOM-138-SEMARNAT/SS-2003 for HFH from industrial-use soil (6000 mg/kg of soil) with an 85% removal at two months. It must be noted that this biopile had a passive aeration system through the vertical and horizontal installation of grooved tubes, placing cones to catch the wind from the dominant direction on the upper ends of the tubes.

The following graphs depict the behavior of 3 of the 14 constructed biopiles. These three were chosen, considering that they are the most representative ones (Figures 2–4).

**Microbiological results**

A microbiological assessment was performed at the start of the project, consisting in bacteria counting tests based on the plate-counting method indicated by the NOM-092-SSA1-1994 [20]. These tests were applied to representative samples considered as initial samples and collected by the quartering method. 

![Figure 2: Behavior of biopile 2 in the studied time.](image-url)
limited by a clayed-lime stratum, which gives the aquifer characteristics of low oxygen transfer, hence the number of aerobic bacteria, which are the ones considered for the aforementioned tests, was quite low.

During the bioremediation treatment, the number of CFU found increased gradually until reaching an optimal amount for the specific conditions of the treatment, which favored degradation of the hydrocarbons present in the soil. In average, the number of CFU increased in two to three orders of magnitude in all biopiles. Some authors, who performed an experiment to compare conventional aeration with a new system proposed by them, found that the heterotrophic microbial population, after 15 days of treatment, increased one order of magnitude, decreasing afterwards, which is a common pattern of bacterial growth [22].

The microbiological analysis was complemented with the identification of bacteria, of which eight species could be isolated and identified. These species are considered to be hydrocarbons degrading bacteria, and were: Vibrio metchnikovii, Micrococcus kirstiae, Pseudomonas luteola, Bacillus brevis, Bacillus megaterium, Bacillus licheniformes, Pseudomonas aeruginosa, Flavimonas oryzaehabitans. Identification was achieved with BBL and API kits.

Respirometry values in the biopiles, in which CO2 generation could be observed and is indicative of biodegradation, were in the 2.46 to 4.0 mg CO2/kg of soil per day range, which, according to some authors [23], corresponds to a low degradation rate, but which sufficed to attain results with HFH, complying with the established normativity. However, other authors report lower respirometry values and consider them to represent an adequate degradation activity. An author reported a degradation rate from 32 µg CO2/g soil to 150 µg CO2/g soil, after 61 to 161 days of incubation in five different treatments [24].

The factors that influence biodegradation (pH, energy source, soil temperature, water and nutrients content, aeration) are discussed in the following.

- The basic energy source in this project was constituted by hydrocarbons, since the organic matter content of the soil was very low as to constitute an important energy source.
- The pH of the soil (7.16) was within the indicated range (6.5-7.5).
- Temperature of soil in the biopiles. This temperature increased once the nutrients were added. During pretreatment, the soil temperature was of 18°C, increasing in the biopiles to 22°C, in average.
- Aeration. Air supply was calculated to obtain the required oxygen of 2 kg O2/kg of HFH. However, the possibility that anaerobic degradation could have occurred in the center of the biopile cannot be discarded because of the presence of CO2 generated in the aerobic zone.
- Nutrients (nitrogen and phosphorus) supplemented through the commercial fertilizers were an important substrate source to facilitate degradation. For 16 000 mg/kg carbon content, nitrogen was added at 1150 mg/kg since the concentration of the site’s soil was 450 mg/kg, and the added phosphorus content was of 157 mg/kg since the original P concentration was of 3 mg/kg. With this, the C:N:P ratio of 100:10:1 was attained.

**Conclusion**

The biopiles remediation method applied to the Marine Terminal “Dos Bocas”, in the state of Tabasco, resulted adequate to remove hydrocarbons in 27 400 m3 of soil, since the final HFH concentrations were below the maximum permissible limits established by the NOM-
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