Evaluation of Biological Treatability of Soil Contaminated with Manufactured Gas Plant Waste

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ABSTRACT

The biological treatability of subsurface soil contaminated with manufactured gas plant (MGP) waste was evaluated. Mineralization assays incorporating ¹⁴C-phenanthrene were used to evaluate the biotransformation potential of indigenous microorganisms at the site. Multi-phase laboratory microcosms were used to evaluate the interphase transfer potential and chemical mass distribution of phenanthrene. The Microtox™ bioassay was used to evaluate detoxification trends at the site. Mineralization results indicated that indigenous microorganisms at the site were capable of transforming phenanthrene, a component of coal-tar creosote. Results also indicated that spiked ¹⁴C-phenanthrene mineralization was influenced by nutrient addition and by the amount of contamination. The chemical mass distribution of ¹⁴C-phenanthrene indicated that volatilization may be an important transport mechanism for chemicals residing in, or migrating to the vadose zone of soil. Following removal of the coal-tar waste source at the site, the toxicity of water soluble extracts of the site soil decreased to a non-toxic response based upon Microtox™ assay results. Parent compound concentrations at the site also decreased with time subsequent to source removal. Results of this study indicate that natural in situ bioremediation may be an important treatment process at a former manufactured gas plant waste site in New York.

INTRODUCTION

The application of wood-preserving, petroleum, and manufactured gas plant (MGP) wastes on soil frequently results in contamination of soil and aquifer material with polycyclic aromatic hydrocarbons (PAHs)[1]. Several PAHs have been identified as carcinogenic and therefore present potential health-related risks with chronic exposure [2]. The biodegradation of phenanthrene (a model three-ring PAH) in liquid cultures by single bacterial species has been demonstrated [3-9]; however, less information is available regarding the biodegradation of phenanthrene or other PAHs in complex coal-tar contaminated aquifers.

The biological treatability of coal-tar contaminated soil and aquifer material is influenced by the extent of contamination. The extent of soil and aquifer contamination requires an evaluation of the aquifer solid phase with regard to hazardous parent chemicals and the natural metabolic potential of indigenous microorganisms to transform target hazardous compounds. The National Research Council (1993), in their recent publication "In situ Bioremediation; When Does it Work?", outlined a strategy combining the collection of field and laboratory data to evaluate the use of in situ bioremediation. The measurement of parent compound disappearance in the field, and the demonstration of the metabolic potential of indigenous microorganisms at a site to transform target parent compounds through laboratory experiments, are primary objectives for evaluating the efficacy of in situ bioremediation [10].
To assess in situ PAH biotransformation in a coal-tar contaminated soil, experiments were conducted utilizing $^{14}$C-phenanthrene. A mineralization assay, developed at Cornell University, was employed using a coal-tar contaminated soil spiked with radiolabeled $^{14}$C-phenanthrene at 22-24°C. This study was conducted to evaluate the biodegradability and transformation potential of phenanthrene in contaminated soil from the saturated and unsaturated zone of a manufactured gas plant waste (MGP) site. Additionally, a study was carried out to evaluate the chemical mass balance distribution of $^{14}$C-C02 resulting from the biotransformation of $^{14}$C-phenanthrene. The toxicity of aqueous extracts of soil from the site was also determined.

METHODS AND MATERIALS

Field Samples

The MGP waste site for this study (EBOS site 24) has been described elsewhere [10]. Soil cores were obtained from EBOS site 24 subsequent to source removal. The source material at the site was removed in the summer of 1991 [11]. Three locations at the site were used for repeated core sampling. A first core was taken in the original source area and designated Station #1. A second core was taken outside the contaminated plume and designated Station #2. This core served as the control sample. A third core was taken down-gradient from the original source material (approx. 200 m) [10] within the contaminated plume and designated Station #19.

Mineralization Assay

A core sample from each station was obtained aseptically [12] and sampled at three depths: (1) 7.5-9.5 ft below ground surface, (water table interface, WT) (2) 9.5-13.5 ft below ground surface, (saturated zone 1, S1) and (3) 12.5-19.5 ft below ground surface (saturated zone 2, S2). Three grams of soil from each depth in each core were aseptically weighed into 25 ml septum vials (Pierce, Rockford, Illinois). Triplicate vials plus one poisoned control were used for each sediment. Approximately 11000 DPMs of $^{9-14}$C-phenanthrene (Sigma Corp., St. Louis, MO) and 0.11 μg of unlabeled phenanthrene (saturated) were added to each vial. To the poisoned controls, 0.5 ml of a 2% solution of HgCl$_2$ was added. Approximately 0.5 ml of 0.5 N NaOH was used as a $^{14}$C-C02 trapping solution and was contained in 15x45 mm shell vials (VWR Scientific, Greenbelt, MD) placed inside the 25 ml septum vials. The inner vials were elevated above the soil by a glass marble. The base trapping solution was removed with a glass syringe (Hamilton Company, Reno, Nevada) from the 25 ml septum vials on days 1, 2, 5, 8 and 19 and analyzed for evolved $^{14}$C-C02 using a Beckman (Model LS 6000 SE) liquid scintillation counter (Beckman Instruments, Redmond, Washington). Fresh base trapping solution was added subsequent to each sampling period. Incubations were carried out at 20-22°C in the dark.

Chemical Mass Distribution

The biotransformation and chemical mass balance distribution of added $^{14}$C-phenanthrene in site soil was evaluated using a Biometer flask [13] system. Approximately 50 g of soil from station #2 (depth 2 ft.) were added to seven separate biometer flasks. The moisture content of the soil was determined to be approximately 15% dry wt. Nutrient analysis of the soil was carried out by Cornell University in 1989 [10] and showed available NH$_3$ and NO$_3$ in the unsaturated zone to be less than 9 mg/Kg, combined. Each of the seven microcosms was spiked with 20 μCi/Kg (wet weight) of $^{9-14}$C-phenanthrene (Sigma Corp., St. Louis, MO) and 200 mg/Kg of unlabeled phenanthrene. One microcosm served as a control and was poisoned by saturating the soil with a 2% HgCl$_2$ solution. Evolved $^{14}$C-C02 was trapped in 10 ml of 0.1 N KOH held in the side arm of each biometer flask. A 60 ml syringe was used to remove and replenish the base trapping solution from the side arm of the biometer flask. Aeration of each biometer microcosm was achieved by drawing 150 ml of fresh air through the system at each sampling period with the syringe. The fresh air was passed through an ascarite column, mounted on top the biometer flask, to remove CO$_2$ prior to aeration. This aeration
procedure also enhanced the entrapment of any volatile $^{14}$C within the polyurethane plug for subsequent analysis.

Analysis of evolved $^{14}$C-$\text{CO}_2$ was carried out by evacuating the 10 ml trapping solution and analyzing 1 ml portions (diluted to 10 ml with scintillation cocktail) using a Beckman (Model LS 6000 SE) liquid scintillation counter (Beckman Instruments, Redmond, Washington). Fresh $\text{CO}_2$ trapping solution was replaced in the side arm of the biometer flask after each sampling. At day 17, three biometer flask microcosms (S1, S2, S3 Table 1) were sacrificed to determine the distribution of $^{14}$C-carbon among phases within the biometer system. The chemical mass distribution was determined by measuring: 1) cumulative $^{14}$C-$\text{CO}_2$ evolved, 2) volatilized $^{14}$C-carbon in the polyurethane foam plug, 3) solvent extractable $^{14}$C-carbon in the soil, and 4) non-solvent extractable $^{14}$C-carbon in the soil. The remaining four microcosms, including the poisoned control, were sacrificed at day 65. A summary of the experimental matrix is given in Table 1.

Volatilized $^{14}$C-carbon was Soxhlet extracted from the polyurethane plug associated with each biometer flask, with approximately 250 ml of methylenechloride/acetone (1:1 vol:vol) for 16-24 hours. The solvent extract was then condensed to approximately 1.5 ml and brought to a final volume of 10 ml in acetonitrile. A portion (1 ml) from each organic plug extract was mixed with approximately 10 ml of scintillation cocktail and counted by liquid scintillation.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Abbreviation</th>
<th>Definition</th>
<th>Sacrificed on Day</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>S1</td>
<td>Microcosm 1</td>
<td>17</td>
</tr>
<tr>
<td>2</td>
<td>S2</td>
<td>Microcosm 2</td>
<td>17</td>
</tr>
<tr>
<td>3</td>
<td>S3</td>
<td>Microcosm 3</td>
<td>17</td>
</tr>
<tr>
<td>4</td>
<td>UN</td>
<td>Unknown</td>
<td>65</td>
</tr>
<tr>
<td>5</td>
<td>NP1</td>
<td>Microcosm with nitrogen and phosphorous addition at day 39: (20 mg-N/Kg, 1.0 mg-P/Kg)</td>
<td>65</td>
</tr>
<tr>
<td>6</td>
<td>NP2</td>
<td>Microcosm with nitrogen and phosphorous addition at day 39: (14 mg-N/Kg, 1.0 mg-P/Kg)</td>
<td>65</td>
</tr>
<tr>
<td>7</td>
<td>PC</td>
<td>Poisoned Control</td>
<td>65</td>
</tr>
</tbody>
</table>

The solvent extractable $^{14}$C-carbon was determined by extracting the soil phase of each biometer flask with methylenechloride/acetone (1:1 vol:vol) for 6 minutes using a sonic agitation probe (Heat Systems, Inc., Farmingdale, NY). The solvent extract was then condensed to approximately 1.5 ml and brought to a final volume of 10 ml in acetonitrile. A portion (1 ml) from each solvent extract was mixed with approximately 10 ml of scintillation cocktail and counted by liquid scintillation.

Non-solvent extractable residues of $^{14}$C-carbon were analyzed by combusting (in triplicate) 1.5 g portions of soil from each biometer flask microcosm. Samples were combusted at 900°C using a Biological Oxidizer OX600 (R.J. Harvey Instrument Corp., Hillsdale, New Jersey). Liberated $^{14}$C-carbon was trapped in a 20 ml solution containing, on a volume to volume basis, 50% ready gel (Beckman Ready Gel, Beckman Inst., Redmond, Washington), 40% reagent grade methanol, and 10% reagent grade monoethanolamine. The radioactivity trapped in the total volume of trapping solution was counted by liquid scintillation.

Toxicity

Toxicity trends associated with the extent of contamination at the site were determined by: 1) evaluating the mineralization potential of indigenous microorganisms in highly contaminated soil and 2) performing Microtox™ analysis of aqueous soil extracts from soil core material from station #19 at two points in time (1991 and 1993) subsequent to source removal.
Mineralization activity was evaluated in four additional microcosms containing contaminated soil from station #19. A soil core taken 1991 revealed a distinct zone of highly contaminated soil from 4 to 8 feet below the ground surface. Soil at this depth was darker in color compared to soil either above or below this level. The zone of high contamination contained phenanthrene at a concentration of approximately 300 mg/kg, compared to 3 mg/kg at a depth of approximately 10 feet [14]. Approximately 50 g of soil was taken in duplicate from within the highly contaminated zone, and below in the lightly contaminated zone at a depth of approximately 10 feet. Each microcosm was spiked with 20 \( \mu Ci/\text{Kg} \) (wet weight) of 9-\(^{14}\text{C}\)-phenanthrene (Sigma Corp., St. Louis, MO) and incubated at 30°C. Evolved \(^{14}\text{C}\)-CO\(_2\) was trapped in 10 ml of 0.1 N KOH held in the side arm of each biometer flask. Mineralization was evaluated as described for the chemical mass distribution study, however, the distribution of the \(^{14}\text{C}\)-phenanthrene within these four microcosms was not determined.

The Microtox™ bioassay was used to measure the extent of detoxification of contaminated core material from within the plume which had occurred after source removal. The Microtox bioassay utilizes the marine organism \textit{Photobacterium phosphoreum} as a test organism. The Microtox measures the reduction in light output of test organisms when challenged by varying concentrations of aqueous samples as an indication of the degree of toxicity of a particular sample. Approximately 25 g of soil, taken from 4-8 ft depth at Station #19, was extracted with 100 ml of distilled-deionized water for 16-24 hr in duplicate on a rotary tumbler at 30 rpm. The samples were allowed to settle for twenty-four hours prior to analysis. The effective concentration of aqueous sample causing a reduction in light output by 50% is called the EC 50. The lower the EC 50 value, the higher the degree of toxicity or inhibition caused by the organic constituents within the aqueous extract. EC 50 values greater than 100 are considered non-toxic. EC 50 values were determined for soil from station #19 taken in 1991 and 1993.

**RESULTS AND DISCUSSION**

**Mineralization Assay**

Cumulative \(^{14}\text{C}\)-CO\(_2\) production curves for the mineralization of \(^{14}\text{C}\)-phenanthrene in site soil from each station and depth are presented in Figures 1-3. Mineralization of \(^{14}\text{C}\)-phenanthrene indicated the potential for the formation of intermediate or biotransformation products by the indigenous microbial population in coal tar contaminated soil. The lack of an observed lag period indicates that aquifer material from each bore hole appears to have microorganisms that are capable of biodegrading PAH. The low cumulative evolved \(^{14}\text{CO}_2\) (<3.5% for replicates 1-3) from Station #2, saturated zone 2, may have resulted from oxygen or nutrient limitations or the population of microorganisms being unable to adapt and transform phenanthrene.

Cumulative \(^{14}\text{C}\)-CO\(_2\) at each station depth was evaluated by computing the mean and standard deviation of triplicate measurements. For each mean and standard deviation, evolved \(^{14}\text{C}\)-CO\(_2\) in poisoned controls are shown in parenthesis. Mineralization of phenanthrene in sediments from the restored source area, Station #1 (Figure 1), did not vary significantly with depth (Figure 1 a, b, c). Mineralization at the water table interface (8.5 ft), saturated zone 1 (14.5 ft), and saturated zone 2 (19.0 ft) was observed to be 32 \(\pm 0.5\%\) (4.9%), 29.9 \(\pm 7.7\%\) (5.1%), and 33.4 \(\pm 1.2\%\) (3.3%) respectively.

Mineralization of phenanthrene in sediments from Station #2 (Figure 2) appears to vary with depth. Mineralization of 24.5 \(\pm 6.2\%\) (8.1%) was observed at the water table interface depth of 7.5 ft. Mineralization of 16.7 \(\pm 0.8\%\) (14.2%) was observed at the saturated zone 1 depth of 13.5 ft. At the lowest depth of Station #2, 17.8 ft, mineralization of 2.6 \(\pm 0.7\%\) (3.0%) was observed.

Mineralization of phenanthrene in sediments from within the contaminated plume area, Station #19 (Figure 3), was observed to decrease to the lowest with depth. Mineralization at the water table interface (9.7 ft), saturated zone 1 (15.5 ft), and saturated zone 2 (19.5 ft) was observed to be 28.9 \(\pm 2.5\%\) (5.0%), 27.7 \(\pm 10.2\%\) (7.0%), and 15.9 \(\pm 2.5\%\) (11.1%) respectively.

The mineralization activity of each core and depth was evaluated using a first order kinetic model. The rates of mineralization for each core and depth are presented in Table 2. The mineralization activity, evidenced by \(^{14}\text{CO}_2\) production, was found in general, to decrease with depth. The distribution of mineralization activity with depth has also been supported by other
researchers working jointly on the same site [10]. Madsen et al., (1991) observed the decline in microscopic and viable counts of bacteria with depth below the water table [10], which may account for the observed decrease in mineralization rate with depth.

![Graph](a)

![Graph](b)

![Graph](c)

**FIGURE 1**
Evolved $^{14}$CO$_2$ during partial mineralization of $^{14}$C-phenanthrene of Station #1 sediments at three depths; a) water table, depth 8.5 ft, b) saturated zone 1, depth 14.5 ft, and c) saturated zone 2, depth 19.0 ft.
FIGURE 2
Evolved $^{14}$CO$_2$ during partial mineralization of $^{14}$C-phenanthrene of Station #2 sediments at three depths; a) water table, depth 7.5 ft, b) saturated zone 1, depth 13.5 ft, and c) saturated zone 2, depth 17.8 ft.
Evolved $^{14}$C$_2$O$_2$ during partial mineralization of $^{14}$C-phenanthrene of Station #19 sediments at three depths; a) water table, depth 9.7 ft, b) saturated zone 1, depth 15.5 ft, and c) saturated zone 2, depth 19.5 ft.
TABLE 2
First order mineralization rates of $^{14}$C-phenanthrene for each core and depth (%/day) with upper and lower 95% confidence intervals in parenthesis.

<table>
<thead>
<tr>
<th>Depth</th>
<th>Station #1</th>
<th>Station #2</th>
<th>Station #19</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water Table (7.5-9.5 ft)</td>
<td>1.5 (0.004, 3.0)</td>
<td>1.0 (0.1, 2.0)</td>
<td>1.3 (0.4, 2.1)</td>
</tr>
<tr>
<td>Saturated zone 1 (9.5-13.5 ft)</td>
<td>0.9 (.009, 1.8)</td>
<td>0.5 (0.2, 1.2)</td>
<td>1.1 (0.1, 2.4)</td>
</tr>
<tr>
<td>Saturated zone 2 (12.5-19.5 ft)</td>
<td>1.4 (0.2, 2.5)</td>
<td>0.05 (0.01, 0.1)</td>
<td>0.7 (0.1, 1.3)</td>
</tr>
</tbody>
</table>

Chemical Mass Distribution

Biodegradation is often assessed by evaluating the disappearance of target hazardous compounds [3, 15]. However, evaluation of the interphase transfer rates in a multi-phase subsurface environment can provide information for assessing other important mechanisms for loss of parent compounds [16]. A chemical mass distribution was carried out on seven microcosms, containing 50 grams each of soil, from the unsaturated zone (Station #2-2 ft depth) spiked with 9-$^{14}$C-phenanthrene. Less than 0.15% mineralization of the added phenanthrene was observed through 17 days of incubation, for all samples, at 30°C (Figure 4 a, b).

![Graph](a)

![Graph](b)

FIGURE 4
Evolved $^{14}$C-CO$_2$ versus time for EBOS Site 24 (Control) soil spiked with 20 μCi /kg (9)-$^{14}$C-phenanthrene through a) 17 days and b) 65 days.
At 17 days, three microcosms were sacrificed to evaluate the distribution of $^{14}$C-carbon. The mass balance results are presented in Table 3. The majority (approx. 57%) of the added label was observed in the solvent extractable phase while approximately 28% of the added radiolabeled carbon was found in the organic plug extract. The remaining fraction obtained by combusting the soil, was found to contain 5.5% of the total radiolabeled carbon. At day 65, NP1, NP2, PC and UN microcosms were sacrificed to obtain a mass distribution of $^{14}$C-carbon. The results of the mass distribution are presented in Table 3.

Volatilized $^{14}$C-carbon for microcosms receiving added nutrients was observed to be 51-56%. The organic plug extracts of the poisoned control and unamended microcosm contained 56-53% $^{14}$C, respectively. Volatilization of $^{14}$C-carbon increased from over 25% (S1-S3) at day 17, to over 50% (PC, NP1, NP2 and UN) at day 65. Therefore, volatilization appears to play a dominating role in the distribution of the $^{14}$C-carbon within all microcosms at 30°C.

In addition to volatilization, sorption of target compounds to soil material influences the fate of hydrophobic organic compounds in subsurface environments [17]. In this study, sorption was assessed by quantifying $^{14}$C-carbon remaining on the soil (soil combustion) subsequent to solvent extraction. $^{14}$C-carbon bound to the soil accounted for less than 7% of the total radiolabeled carbon added in all samples, with the exception of NP1, where 11% of the radiolabeled carbon was bound to the soil. The sorption of organic compounds to soil can affect the rate of biodegradation [18, 19]. Microbial uptake rates can be limited by slow-desorption since only compounds in the non-sorbed state are typically subject to microbial attack [20].

The rate limiting step in the microbial uptake and subsequent mineralization of phenanthrene in this study, appears to be N and P availability. At day 39, nitrogen and phosphorus were added to microcosm NP1 and NP2, resulting in partial mineralization of 9-14C-phenanthrene (20%-6%, Figure 4 b).

Toxicity Evaluation

Mineralization of $^{14}$C-phenanthrene was also used to evaluate the toxicity of lightly and heavily contaminated soil. Soil from Station #19 containing a distinct zone of high contamination approximately 4 to 8 feet below the ground surface was evaluated for the mineralization potential of indigenous microorganisms. Lightly contaminated soil amended with nutrients resulted in cumulative $%^{14}$C-CO$_2$ production in the range of 25-30% after 5 days of incubation at 30°C. Results are presented in Figure 5. The highly contaminated soil microcosm yielded no $%^{14}$C-CO$_2$ production. Inhibition of mineralization in the highly contaminated soil is attributed to a toxic effect of creosote on the soil microbial population.

The Microtox assay was used to evaluate changes in toxicity with time of core material from the site. EC50 values were determined for core material at Station #19 taken at two different sampling periods in 1991 and 1993, respectively. The results of the Microtox analysis are given in Table 4. The aqueous extracts of the soil core material from 1991 samples yielded a toxic response to the Microtox organisms (EC50 = 17.5). Analyzing aqueous extracts from soil core material from the same location in 1993 resulted in a non-toxic response to the Microtox$^{\text{TM}}$ organisms. The observed trend towards detoxification may have been influenced by the removal of the source material in 1991. The removal of the source material was followed by an observed decrease in plume concentrations of naphthalene and phenanthrene [21]. Decreases in toxicity are probably caused by the removal of

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**TABLE 3**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Time (Days)</th>
<th>Organic Plug Extract</th>
<th>Solvent Extract</th>
<th>Soil Combustion</th>
<th>$^{14}$C-CO$_2$</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>17</td>
<td>30.68</td>
<td>55.60</td>
<td>4.98</td>
<td>0.098</td>
<td>91.35</td>
</tr>
<tr>
<td>S2</td>
<td>17</td>
<td>26.45</td>
<td>58.41</td>
<td>6.41</td>
<td>0.097</td>
<td>91.37</td>
</tr>
<tr>
<td>S3</td>
<td>17</td>
<td>28.84</td>
<td>58.17</td>
<td>5.14</td>
<td>0.099</td>
<td>92.25</td>
</tr>
<tr>
<td>UN</td>
<td>65</td>
<td>53.18</td>
<td>32.94</td>
<td>3.91</td>
<td>0.13</td>
<td>90.16</td>
</tr>
<tr>
<td>NP1</td>
<td>65</td>
<td>51.94</td>
<td>4.41</td>
<td>11.02</td>
<td>20.45</td>
<td>87.82</td>
</tr>
<tr>
<td>NP2</td>
<td>65</td>
<td>56.32</td>
<td>12.05</td>
<td>5.77</td>
<td>6.90</td>
<td>81.04</td>
</tr>
<tr>
<td>PC</td>
<td>65</td>
<td>56.29</td>
<td>14.57</td>
<td>4.39</td>
<td>0.06</td>
<td>75.31</td>
</tr>
</tbody>
</table>
source material entering the plume and the occurrence of in situ biodegradation. Dilution through groundwater flow and recharge may have also influenced toxicity trends at the site.

![Graph showing %14C-CO2 Recovery over time](image)

FIGURE 5

Evolved $^{14}$CO$_2$ from laboratory microcosms, spiked with $^{14}$C-phenanthrene, containing heavily contaminated and lightly contaminated soil from Station #19.

**TABLE 4**

Microtox EC50 results from aqueous extracts of soil within the contaminated plume area at Station #19.

<table>
<thead>
<tr>
<th>SAMPLE</th>
<th>EC50</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Station #19 (1991)</td>
<td>17.5</td>
<td>6.5</td>
</tr>
<tr>
<td>Station #19 (1993)</td>
<td>NT</td>
<td>6.5</td>
</tr>
<tr>
<td>Station #2 (1993)</td>
<td>NT</td>
<td>6.7</td>
</tr>
</tbody>
</table>

NT-non toxic (EC50 value greater than 100)

CONCLUSIONS

Field and laboratory results from this research provide evidence that natural in situ bioremediation is an important treatment process at a former manufactured gas plant waste site in New York. Indigenous soil microorganisms at the site exhibited the potential to transform phenanthrene, a known component of coal tar creosote. The production of $^{14}$C-CO$_2$ from $^{14}$C-labeled phenanthrene indicated that metabolic intermediates may be produced through the oxidation of phenanthrene. This research indicated that microorganisms in both contaminated and uncontaminated soil are capable of transforming phenanthrene in the site soil.

Chemical mass distribution studies using laboratory microcosms indicated that volatilization and nutrient availability may play important roles in the biotreatment of contaminated soil at the site. Reliance on disappearance of parent compounds as an estimate of biodegradation may over-estimate transformation due to natural biological processes. Interphase transfer potential of chemicals in a multiphase subsurface environment should be evaluated.

Microbial processes responsible for biodegradation were inhibited in highly contaminated soil containing coal-tar creosote. Engineering approaches such as source removal can enhance natural bioremediation processes by stopping the introduction of toxic chemicals at a site. Biotransformation products may be more toxic or less toxic than target parent compounds. Examining aqueous extracts of soil core material using the Microtox™ assay provided a simple field measure of toxicity trends at the site and indicated decreasing toxicity with time following source removal.
Monitoring parent compound concentrations and toxicity trends at a site are field measurements useful in evaluating the efficacy of in situ bioremediation. Concurrently, laboratory experiments may be employed to evaluate: (1) the potential of indigenous microorganisms to transform target compounds such as PAH, (2) the influence of nutrients, and (3) the interphase transfer potential of target compounds and transformation products. These field measurements and laboratory assays provide information useful for site characterization and are congruent with the National Research Council's strategy for evaluation of in situ biotreatment of contaminated soils.

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